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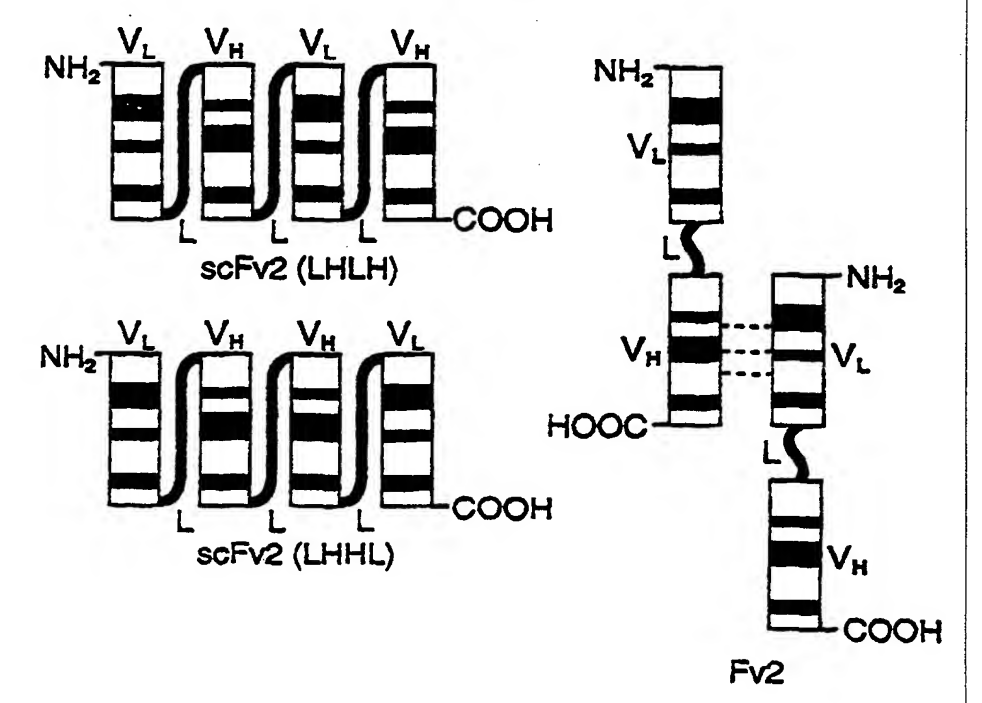
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(54) Title: MULTIVALENT SINGLE CHAIN ANTIBODIES

#### (57) Abstract

The present invention discloses multivalent single chain antibodies which have two or more biologically active antigen binding sites. The multivalent single chain antibodies are formed by using a peptide linker to covalently link two or more single chain antibodies, each single chain antibody having a variable light domain linked to a variable heavy chain domain by a peptide linker.

### Schematic Representation Of Covalently & Non-Covalently Linked Single Chain Fv Multimers



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#### MULTIVALENT SINGLE CHAIN ANTIBODIES

The present invention relates to single chain multivalent antibodies.

Antibodies are proteins belonging to a group of immunoglobulins elicited by the immune system in response to a specific antigen or substance which the body deems foreign. There are five classes of human antibodies, each class having the same basic structure. The basic structure of an antibody is a tetramer, or a multiple thereof, composed of two identical heterodimers each consisting of a light and a heavy chain. The light chain is composed of one variable (V) and one constant (C) domain, while a heavy chain is composed of one variable and three or more constant domains. The variable domains from both the light and heavy chain, designated V<sub>L</sub> and V<sub>H</sub> respectively, determine the specificity of an immunoglobulin, while the constant (C) domains carry out various effector functions.

Amino acid sequence data indicate that each variable domain comprises three complementarity determining regions (CDR) flanked by four relatively conserved framework regions (FR). The FR are thought to maintain the structural integrity of the variable region domain. The CDR have been assumed to be responsible for the binding specificity of individual antibodies and to account for the diversity of binding of antibodies.

As the basic structure of an antibody contains two heterodimers, antibodies are multivalent molecules. For example, the IgG classes have two identical antigen binding sites, while the pentameric IgM class has 10 identical binding sites.

Monoclonal antibodies having identical genetic parentage and binding specificity have been useful both as diagnostic and therapeutic agents. Monoclonal antibodies are routinely produced by hybridomas generated by fusion of mouse lymphoid cells with an appropriate mouse myeloma cell line according to established procedures. The administration of murine antibodies for in vivo therapy and diagnostics in humans is limited however, due to the human anti-mouse antibody response illicited by the human immune system.

Chimeric antibodies, in which the binding or variable regions of antibodies derived from one species are combined with the constant regions of antibodies derived from a different species, have been produced by recombinant DNA methodology. See, for example, Sahagen et al., *J. Immunol.*, 137:1066-1074 (1986); Sun et al., *Proc. Natl. Acad. Sci. USA*, 82:214-218 (1987); Nishimura et al., *Cancer Res.*, 47:999-1005 (1987); and Lie et al. *Proc Natl. Acad. Sci. USA*, 84:3439-3443 (1987) which disclose chimeric antibodies to tumor-associated antigens. Typically, the variable region of a murine antibody is joined with the constant region of a human antibody. It is expected that as such chimeric antibodies are largely human in composition, they will be substantially less immunogenic than murine antibodies.

Chimeric antibodies still carry the Fc regions which are not necessary for antigen binding, but constitute a major portion of the overall antibody structure which affects its pharmacokinetics. For the use of antibodies in immunotherapy or immunodiagnostics, is it

desirable to have antibody-like molecules which localize and bind to the target tissue rapidly and for the unbound material to quickly clear from the body. Generally, smaller antibody fragments have greater capillary permeability and are more rapidly cleared from the body than whole antibodies.

Since it is the variable regions of light and heavy chains that interact with an antigen, single chain antibody fragments (scFvs) have been created with one  $V_L$  and one  $V_H$ , containing all six CDR's, joined by a peptide linker (U.S. Patent 4,946,778) to create a  $V_L$ -L- $V_H$  polypeptide, wherein the L stands for the peptide linker. A scFv wherein the  $V_L$  and  $V_H$  domains are orientated  $V_H$ -L- $V_L$  is disclosed in U.S. Patent 5,132,405.

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As the scFvs have one binding site as compared to the minimum of two for complete antibodies, the scFvs have reduced avidity as compared to the antibody containing two or more binding sites.

than one binding site to enhance the avidity of the polypeptide, and retain or increase their antigen recognition properties. In addition, it would be beneficial to obtain multivalent scFvs which are bispecific to allow for recognition of different epitopes on the target tissue, to allow for antibody-based recruitment of other immune effector functions, or allow antibody capture of a therapeutic or diagnostic moiety.

It has been found that single chain antibody fragments, each having one  $V_H$  and one  $V_L$  domain covalently linked by a first peptide linker, can be covalently linked by a second peptide linker to form a multivalent single chain antibody which maintains the binding affinity of a whole antibody. In one embodiment, the present invention is a multivalent single chain antibody having affinity for an antigen wherein the multivalent single chain antibody comprises two or more light chain variable domains and two or more heavy chain variable domains; wherein, each variable domain is linked to at least one other variable domain.

In another embodiment, the present invention is a multivalent single chain antibody which comprises two or more single chain antibody fragments, each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:

- (a) a first polypeptide comprising a light chain variable domain;
- (b) a second polypeptide comprising a heavy chain variable domain; and
- (c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.

In another embodiment, the invention provides a DNA sequence which codes for a multivalent single chain antibody, the multivalent single chain antibody comprising two or more single chain antibody fragments, each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:

(a) a first polypeptide comprising a light chain variable domain;

(b) a second polypeptide comprising a heavy chain variable domain; and

(c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.

The multivalent single chain antibodies allow for the construction of an antibody fragment which has the specificity and avidity of a whole antibody but are smaller in size allowing for more rapid capillary permeability. Multivalent single chain antibodies also allow for the construction of a multivalent single chain antibody wherein the binding sites can be two different antigenic determinants.

#### BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 illustrates covalently linked single chain antibodies having the configuration  $V_L$ -L- $V_H$ -L- $V_H$  (LHLH) and  $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_L$  (LHHL) and a noncovalently linked Fv single chain antibody (Fv2).

Figure 2 illustrates the nucleotide sequence of CC49 V<sub>L</sub>.

Figure 3 illustrates the amino acid sequence of CC49 V<sub>L</sub>.

Figure 4 illustrates the nucleotide sequence of CC49 V<sub>H</sub>.

Figure 5 illustrates the amino acid sequence of CC49 V<sub>H</sub>.

Figure 6 illustrates the nucleotide sequence and amino acid sequence of the CC49 single chain antibody LHLH in p49LHLH.

Figure 7 illustrates the nucleotide sequence and amino acid sequence of the CC49  $_{20}$  single antibody LHHL in p49LHHL.

Figure 8 illustrates construction of plasmids pSL301 T and pSL301 HT.

Figure 9 illustrates construction of plasmid p49LHHL.

Figure 10 illustrates construction of plasmid p49LHLH.

Figure 11 illustrates the results of a competition assay using CC49 IgG, CC49 scFv2, and CC49 scFv using biotinylated CC49 IgG as competitor.

The entire teaching of all references cited herein are hereby incorporated by reference.

Nucleic acids, amino acids, peptides, protective groups, active groups and such, when abbreviated, are abbreviated according to the IUPAC IUB (Commission on Biological Nomenclature) or the practice in the fields concerned.

The term "single chain antibody fragment" (scFv) or "antibody fragment" as used herein means a polypeptide containing a  $V_L$  domain linked to a  $V_H$  domain by a peptide linker (L), represented by  $V_L$ -L- $V_H$ . The order of the  $V_L$  and  $V_H$  domains can be reversed to obtain polypeptides represented as  $V_H$ -L- $V_L$ . "Domain" is a segment of protein that assumes a discrete function, such as antigen binding or antigen recognition.

A "multivalent single chain antibody" means two or more single chain antibody fragments covalently linked by a peptide linker. The antibody fragments can be joined to form bivalent single chain antibodies having the order of the  $V_L$  and  $V_H$  domains as follows:

 $V_L-L-V_H-L-V_L-V_H$ ;  $V_L-L-V_H-L-V_L$ ;  $V_H-L-V_L-V_H-L-V_L$ ; or  $V_H-L-V_L-L-V_L-L-V_H$ . Single chain multivalent antibodies which are trivalent and greater have one or more antibody fragments joined to a bivalent single chain antibody by an additional interpeptide linker. In a preferred embodiment, the number of  $V_L$  and  $V_H$  domains is equivalent.

The present invention also provides for multivalent single chain antibodies which can be designated  $V_H-L-V_L-L-V_L$  or  $V_L-L-V_H-L-V_H$ .

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Covalently linked single chain antibodies having the configuration  $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_H$  (LHLH) and  $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_L$  (LHHL) are illustrated in Figure 1. A noncovalently linked Fv single chain antibody (Fv2) is also illustrated in Figure 1.

The single chain antibody fragments for use in the present invention can be derived from the light and/or heavy chain variable domains of any antibody. Preferably, the light and heavy chain variable domains are specific for the same antigen. The individual antibody fragments which are joined to form a multivalent single chain antibody may be directed against the same antigen or can be directed against different antigens.

To prepare a vector containing the DNA sequence for a single chain multivalent antibody, a source of the genes encoding for these regions is required. The appropriate DNA sequence can be obtained from published sources or can be obtained by standard procedures known in the art. For example, Kabat et al., Sequences of Proteins of Immunological Interest 4th ed., (1991), published by The U.S. Department of Health and Human Services, discloses sequences of most of the antibody variable regions which have been described to date.

When the genetic sequence is unknown, it is generally possible to utilize cDNA sequences obtained from mRNA by reverse transcriptase mediated synthesis as a source of DNA to clone into a vector. For antibodies, the source of mRNA can be obtained from a wide range of hybridomas. See, for example, the catalogue ATCC Cell Lines and Hybridomas, American Type Culture Collection, 20309 Parklawn Drive, Rockville Md., USA (1990). Hybridomas secreting monoclonal antibodies reactive with a wide variety of antigens are listed therein, are available from the collection, and usable in the present invention. These cell lines and others of similar nature can be utilized as a source of mRNA coding for the variable domains or to obtain antibody protein to determine amino acid sequence of the monoclonal antibody itself.

Variable regions of antibodies can also be derived by immunizing an appropriate vertebrate, normally a domestic animal, and most conveniently a mouse. The immunogen will be the antigen of interest, or where a hapten, an antigenic conjugate of the hapten to an antigen such as keyhole limpet hemocyanin (KLH). The immunization may be carried out conventionally with one or more repeated injections of the immunogen into the host mammal, normally at two to three week intervals. Usually, three days after the last challenge, the spleen is removed and dissociated into single cells to be used for cell fusion to provide hybridomas from which mRNA can readily be obtained by standard procedures known in the art.

When an antibody of interest is obtained, and only its amino acid sequence is known, it is possible to reverse translate the sequence.

The  $V_L$  and  $V_H$  domains for use in the present invention are preferably obtained from one of a series of CC antibodies against tumor-associated glycoprotein 72 antigen (TAG-72) disclosed in published PCT Application WO 90/04410 on May 3, 1990, and published PCT Application WO 89/00692 on January 26, 1989. More preferred are the  $V_L$  and  $V_H$  domains from the monoclonal antibody designated CC49 in PCT Publications WO 90/04410 and WO 89/00692. The nucleotide sequence (SEQ ID NO: 1) which codes for the  $V_L$  of CC49 is substantially the same as that given in Figure 1. The amino acid sequence (SEQ ID NO: 2) of the  $V_L$  of CC49 is substantially the same as that given in Figure 2. The nucleotide sequence (SEQ ID NO: 3) which codes for the  $V_H$  of CC49 is substantially the same as that given in Figure 3. The amino acid sequence (SEQ ID NO: 4) for the  $V_H$  of CC49 is substantially the same as that given in Figure 4.

To form the antibody fragments and multivalent single chain antibodies of the present invention, it is necessary to have a suitable peptide linker. Suitable linkers for joining the  $V_H$  and  $V_L$  domains are those which allow the  $V_H$  and  $V_L$  domains to fold into a single polypeptide chain which will have a three dimensional structure very similar to the original structure of a whole antibody and thus maintain the binding specificity of the whole antibody from which antibody fragment is derived. Suitable linkers for linking the scFvs are those which allow the linking of two or more scFvs such that the  $V_H$  and  $V_L$  domains of each immunoglobulin fragment have a three dimensional structure such that each fragment maintains the binding specificity of the whole antibody from which the immunoglobulin fragment is derived. Linkers having the desired properties can be obtained by the method disclosed in U.S. Patent 4,946,778, the disclosure of which is hereby incorporated by reference. From the polypeptide sequences generated by the methods described in the 4,946,778, genetic sequences coding for the polypeptide can be obtained.

Preferably, the peptide linker joining the  $V_H$  and  $V_L$  domains to form a scFv and the peptide linker joining two or more scFvs to form a multivalent single chain antibody have substantially the same amino acid sequence.

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It is also necessary that the linker peptides be attached to the antibody fragments such that the binding of the linker to the individual antibody fragments does not interfere with the binding capacity of the antigen recognition site.

A preferred linker is based on the helical linker designated 205C as disclosed in Pantoliano et al. *Biochem.*, 30, 10117-10125 (1991) but with the first and last amino acids changed because of the codon dictated by the Xho I site at one end and the Hind III site at the other. The amino acid sequence (SEQ ID NO: 5) of the preferred linker is as follows:

The linker is generally 10 to 50 amino acid residues. Preferably, the linker is 10 to 30 amino acid residues. More preferably the linker is 12 to 30 amino acid residues. Most preferred is a linker of 15 to 25 amino acid residues.

Expression vehicles for production of the molecules of the invention include plasmids or other vectors. In general, such vectors contain replicon and control sequences which are derived from species compatible with a host cell. The vector ordinarily carries a replicon site, as well as specific genes which are capable of providing phenotypic selection in transformed cells. For example, *E. coli* is readily transformed using pBR322 [Bolivar et al., *Gene*, 2, 95- (1977), or Sambrook et al., *Molecular Cloning*, Cold Spring Harbor Press, New York, 2nd Ed. (1989)].

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Plasmids suitable for eukaryotic cells may also be used. *S. cerevisiae*, or common baker's yeast, is the most commonly used among eukaryotic microorganisms, although a number of other strains, such as Pichia pastoris, are available. Cultures of cells derived from multicellular organisms such as SP2/0 or Chinese Hamster Ovary (CHO), which are available from the ATCC, may also be used as hosts. Typical of vector plasmids suitable for mammalian cells are pSV2neo and pSV2gpt (ATCC); pSVL and pKSV-10 (Pharmacia), pBPV-1/pML2d (International Biotechnology, Inc.).

The use of prokaryotic and eukaryotic viral expression vectors to express the genes for polypeptides of the present invention is also contemplated.

It is preferred that the expression vectors and the inserts which code for the single chain multivalent antibodies have compatible restriction sites at the insertion junctions and that those restriction sites are unique to the areas of insertion. Both vector and insert are treated with restriction endonucleases and then ligated by any of a variety of methods such as those described in Sambrook et al., supra.

Preferred genetic constructions of vectors for production of single chain multivalent antibodies of the present invention are those which contain a constitutively active transcriptional promoter, a region encoding signal peptide which will direct synthesis/secretion of the nascent single chain polypeptide out of the cell. Preferably, the expression rate is commensurate with the transport, folding and assembly steps to avoid accumulation of the polypeptide as insoluble material. In addition to the replicon and control sequences, additional elements may also be needed for optimal synthesis of single chain polypeptide. These elements may include splice signals, as well as transcription promoter, enhancers, and termination signals. Furthermore, additional genes and their products may be required to facilitate assembly and folding (chaperones).

Vectors which are commercially available can easily be altered to meet the above criteria for a vector. Such alterations are easily performed by those of ordinary skill in the art in light of the available literature and the teachings herein.

Additionally, it is preferred that the cloning vector contain a selectable marker, such as a drug resistance marker or other marker which causes expression of a selectable trait by the host cell. "Host cell" refers to cells which can be recombinantly transformed with vectors constructed using recombinant DNA techniques. A drug resistance or other selectable marker is intended in part to facilitate in the selection of transformants. Additionally, the presence of a selectable marker, such as a drug resistance marker, may be of use in keeping contaminating microorganisms from multiplying in the culture medium. In this embodiment, such a pure culture of the transformed host cell would be obtained by culturing the cells under conditions which require the induced phenotype for survival.

Recovery and purification of the present invention can be accomplished using standard techniques known in the art. For example, if they are secreted into the culture medium, the single chain multivalent antibodies can be concentrated by ultrafiltration. When the polypeptides are transported to the periplasmic space of a host cell, purification can be accomplished by osmotically shocking the cells, and proceeding with ultrafiltration, antigen affinity column chromatography or column chromatography using ion exchange chromatography and gel filtration. Polypeptides which are insoluble and present as refractile bodies, also called inclusion bodies, can be purified by lysis of the cells, repeated centrifugation and washing to isolate the inclusion bodies, solubilization, such as with guanidine-HCl, and refolding followed by purification of the biologically active molecules.

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The activity of single chain multivalent antibodies can be measured by standard assays known in the art, for example competition assays, enzyme-linked immunosorbant assay (ELISA), and radioimmunoassay (RIA).

The multivalent single chain antibodies of the present invention provide unique benefits for use in diagnostics and therapeutics. The use of multivalent single chain antibodies afford a number of advantages over the use of larger fragments or entire antibody molecules. They reach their target tissue more rapidly, and are cleared more quickly from the body.

For diagnostic and/or therapeutic uses, the multivalent single chain antibodies can be constructed such that one or more antibody fragments are directed against a target tissue and one or more antibody fragments are directed against a diagnostic or therapeutic agent.

The invention also concerns pharmaceutical compositions which are particularly advantageous for use in the diagnosis and/or therapy of diseases, such as cancer, where target antigens are often expressed on the surface of cells. For diagnostic and/or therapeutic uses, the multivalent single chain antibodies can be conjugated with an appropriate imaging or therapeutic agent by methods known in the art. The pharmaceutical compositions of the invention are prepared by methods known in the art, e.g., by conventional mixing, dissolving or lyophilizing processes.

The invention will be further clarified by a consideration of the following examples, which are intended to be purely exemplary of the present invention.

-8-

#### **ABBREVIATIONS**

BCIP 5-bromo-4-chloro-3-indoyl phosphate base pair pp (1,3-bis[tris(hydroxymethyl)-methylamino]-Bis-Tris 5 propane) propane bovine serum albumin BSA Complementarity determining region CDR enzyme linked immunosorbent assay ELISA non-covalent single chain Fv dimer Fv2 10 isoelectric focusing IEF kilo base pair Kbp LB Luria-Bertani medium monoclonal antibody Mab 2-(N-Morpholino)ethane sulfonic acid MES 15 molecular weight WW nitro blue tetrazolium chloride NBT Oligo Oligonucleotides polyacrylamide gel PAG polyacrylamide gel electrophoresis PAGE 20 phosphate buffered saline PBS PCR polymerase chain reaction plasmid containing DNA sequence coding for SCFV pSCFV radioimmunoguided surgery RIGS radioimmunotherapy RIT 25 single chain Fv immunoglobulin fragment monomer scFv single chain Fv immunoglobulin fragment dimer scFv2 covalently linked sodium dodecyl sulfate SDS Tris-buffered saline TBS 30 (Tris[hydroxymethyl]aminomethane) Tris Tween-20 wash solution TTBS immunoglobulin heavy chain variable domain  $V_{H}$ immunoglobulin light chain variable domain  $V_L$ 

#### **Antibodies**

CC49: A murine monoclonal antibody specific to the human tumor-associated glycoprotein 72 (TAG-72) deposited as ATCC No. HB9459.

CC49 FAB: An antigen binding portion of CC49 consisting of an intact light chain linked to the N-terminal portion of the heavy chain.

<u>CC49 scFv</u>: Single chain antibody fragment consisting of two variable domains of CC49 antibody joined by a peptide linker.

CC49 Fv2: Two CC49 scFv non-covalently linked to form a dimer. The number after Fv refers to the number of monomer subunits of a given molecule, e.g., CC49 Fv6 refers to the hexamer multimers.

<u>CC49 scFv2</u>: Covalently-linked single chain antibody fragment consisting of two CC49  $V_L$  domains and two  $V_H$  domains joined by three linkers. Six possible combinations for the order of linking the  $V_L(L)$  and the  $V_H(H)$  domains together are: LHLH, LHHL, LLHH, HLLH, and HHLL.

#### 15 Plasmids

pSCFV UHM: Plasmid containing coding sequence for scFv consisting of a CC49 variable light chain and a CC49 variable heavy chain joined by a 25 amino acid linker.

<u>p49LHLH or p49LHHL</u>: Plasmids containing the coding sequence for producing CC49 scFv2 LHLH or LHHL products, respectively.

#### 20 EXAMPLES

#### **General Experimental**

Procedures for molecular cloning are as those described in Sambrook et al.,

Molecular Cloning, Cold Spring Harbor Press, New York, 2nd Ed. (1989) and Ausubel et al.,

Current Protocols in Molecular Biology, John Wiley and Sons, New York (1992), the disclosures of which are hereby incorporated by reference.

All water used throughout was deionized distilled water.

#### Oligonucleotide Synthesis and Purification

All oligonuclotides (oligos) were synthesized on either a Model 380A or a Model 391 DNA Synthesizer from Applied Biosystems (Foster City, CA) using standard β-cyanoethyl phosphoramidites and synthesis columns. Protecting groups on the product were removed by heating in concentrated ammonium hydroxide at 55°C for 6 to 15 hours. The ammonium hydroxide was removed through evaporation and the crude mixtures were resuspended in 30 to 40 μL of sterile water. After electrophoresis on polyacrylamide-urea gels, the oligos were visualized using short wavelength ultraviolet (UV) light. DNA bands were excised from the gel and eluted into 1 mL of 100 mM Tris-HCl, pH 7.4, 500 mM NaCl, 5 mM EDTA over 2 hours at 65°C. Final purification was achieved by applying the DNA to Sep-Pac<sup>TM</sup> C-18 columns (Millipore, Bedford, MA) and eluting the bound oligos with 60 percent methanol. The

solution volume was reduced to approximately 50  $\mu$ L and the DNA concentration was determined by measuring the optical density at 260 nm (OD<sub>260</sub>). Restriction Enzyme Digests

All restriction enzyme digests were performed using Bethesda Research
Laboratories (Gaithersburg, MD), New England Biolabs, Inc. (Beverly, MA) or Boehringer
Mannheim (BM, Indianapolis, IN) enzymes and buffers following the manufacturer's
recommended procedures. Digested products were separated by polyacrylamide gel
electrophoresis (PAGE). The gels were stained with ethidium bromide, the DNA bands were
visualized using long wavelength UV light and the DNA bands were then excised. The gel slices
were placed In dialysis tubing (Union Carbide Corp., Chicago) containing 5 mM Tris, 2.5 mM
acetic acid, 1 mM EDTA, pH 8.0 and eluted using a Max Submarine electrophoresis apparatus
(Hoefer Scientific Instruments, CA). Sample volumes were reduced on a Speed Vac
Concentrator (Savant Instruments, Inc., NY). The DNA was ethanol precipitated and redissolved
in sterile water.

#### 5 Enzyme Linked Immunosorbent Assay (ELISA)

TAG-72 antigen, prepared substantially as described by Johnson et al, Can. Res., 46, 850-857 (1986), was adsorbed onto the wells of a polyvinyl chloride 96 well microtiter plate (Dynatech Laboratories, Inc., Chantilly, VA) by drying overnight. The plate was blocked with 1 percent BSA in PBS for 1 hour at 31°C and then washed 3 times with 200 µL of PBS, 0.05 percent Tween-20. 25 μL of test antibodies and 25 μL of biotinylated CC49 (1/20,000 dilution of a 1 mg/mL solution) were added to the wells and the plate incubated for 30 minutes at 31°C. The relative amounts of TAG-72 bound to the plate, biotinylated CC49, streptavidinalkaline phosphatase, and color development times were determined empirically in order not to have excess of either antigen or biotinylated CC49, yet have enough signal to detect competition by scFv. Positive controls were CC49 at 5 µg/mL and CC49 Fab at 10 µg/mL. Negative controls were 1 percent BSA in PBS and/or concentrated LB. Unbound proteins were washed away. 50 µL of a 1:1000 dilution of streptavidin conjugated with alkaline phosphatase (Southern Biotechnology Associates, Inc., Birmingham, AL) were added and the plate was incubated for 30 minutes at 31°C. The plate was washed 3 more times. 50 µL of a para-nitrophenyl-phosphate solution (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, MD) were added and the color reaction was allowed to develop for a minimum of 20 minutes. The relative amount of scFv2 binding was measured by optical density scanning at 404-450 nm using a microplate reader (Molecular Devices Corporation, Manlo Park, CA). Binding of the scFv2 species resulted in decreased binding of the biotinylated CC49 with a concomitant decrease in color development.

#### SDS-PAGE and Western Biotting

Samples for SDS-PAGE analysis (20  $\mu$ L) were prepared by boiling in a non-reducing sample preparation buffer-Seprasol I (Integrated Separation Systems (ISS), Natick, MA) for

5 minutes and loaded on 10-20 percent gradient polyacrylamide Daiichi Minigels as per the manufacturer's directions (ISS).

Electrophoresis was conducted using a Mini 2-gel apparatus (ISS) at 55 mA per gel at constant current for approximately 75 minutes. Gels were stained in Coomassie Brilliant Blue R-250 (Bio-Rad, Richmond, CA) for at least 1 hour and destained. Molecular weight standards were prestained (Mid Range Kit, Diversified Biotech, Newton Center, MA) and included the following proteins: Phosphorylase b, glutamate dehydrogenase, ovalbumin, lactate dehydrogenase, carbonic amhydrase, B-lactoglobulin and cytochrome C. The corresponding MWs are: 95,500, 55,000, 43,000, 36,000, 29,000, 18,400, and 12,400, respectively.

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When Western analyses were conducted, a duplicate gel was also run. After electrophoresis, one of the gels was equilibrated for 15-20 minutes in anode buffer #1 (0.3 M Tris-HCl pH 10.4). An Immobilon-P PVDF (polyvinylidene dichlorine) membrane (Millipore, Bedford, MA) was treated with methanol for 2 seconds, and immersed in water for 2 minutes. The membrane was then equilibrated in anode buffer #1 for 3 minutes. A Milliblot-SDE apparatus (Millipore) was utilized to transfer proteins in the gel to the membrane. A drop of anode buffer #1 was placed in the middle of the anode electrode surface. A sheet of Whatman 3MM filter paper was soaked in anode buffer #1 and smoothly placed on the electrode surface. Another filter paper soaked in anode buffer #2 (25 mM tris pH 10.4) was placed on top of the first one. A sandwich was made by next adding the wetted PVDF membrane, placing the equilibrated gel on top of this and finally adding a sheet of filter paper soaked in cathode buffer (25mM Tris-HCl, pH 9.4 in 40 mM glycine). Transfer was accomplished in 30 minutes using 250 mA constant current (initial voltage ranged from 8-20 volts).

After blotting, the membrane was rinsed briefly in water and placed in a dish with 20 mL blocking solution (1 percent bovine serum albumin (BSA) (Sigma, St. Louis, MO) in Tris-buffered saline (TBS)). TBS was purchased from Pierce Chemical (Rockford, IL) as a preweighed powder such that when 500 mL water is added, the mixture gives a 25 mM Tris, 0.15 M sodium chloride solution at pH 7.6. The membranes were blocked for a minimum of 1 hour at ambient temperature and then washed 3 times for 5 minutes each using 20 mL 0.5 percent Tween-20 wash solution (TTBS). To prepare the TTBS, 0.5mL of Tween 20 (Sigma) was mixed per liter of TBS. The probe antibody used was 20 mL biotinylated FAID14 solution (10 µg per 20 mL antibody buffer). Antibody buffer was made by adding 1 g BSA per 100 mL of TTBS. After probing for 30-60 minutes at ambient temperature, the membrane was washed 3 times with TTBS, as above.

Next, the membrane was incubated for 30-60 minutes at ambient temperature with 20 mL of a 1:500 dilution in antibody buffer of streptavidin conjugated with alkaline phosphatase (Southern Biotechnology Associates, Birmingham, AL). The wash step was again repeated after this, as above. Prior to the color reaction, membranes were washed for 2 minutes in an alkaline carbonate buffer (20 mL). This buffer is 0.1 M sodium bicarbonate,

1 mM MgCl<sub>2</sub>·H<sub>2</sub>0, pH 9.8. To make up the substrate for alkaline phosphatase, nitroblue tetrazolium (NBT) chloride (50 mg, Sigma) was dissolved in 70 percent dimethylformamide. 5-Bromo-4-chloro-3-indoyl phosphate (BCIP) (25 mg, Sigma) was separately dissolved in 100 percent dimethylformamide. 5-Bromo-4-chloro-3-indoyl phosphate (BCIP) 25 mg, Sigma) was separately dissolved in 100 percent dimethylformamide. These solutions are also commercially available as a Western developing agent sold by Promega. For color development, 120 µL of each were added to the alkaline solution above and allowed to react for 15 minutes before they were washed from the developed membranes with water. Biotinylated FAID14

FAID14 is a murine anti-idiotypic antibody (IgG2a, K isotype) deposited as ATCC No. CRL 10256 directed against CC49. FAID14 was purified using a Nygene Protein A affinity column (Yonkers, NY). The manufacturer's protocol was followed, except that 0.1 M sodium citrate, pH 3.0 was used as the elution buffer. Fractions were neutralized to pH  $\sim$ 7 using 1.0 M Tris-HCl pH 9.0. The biotinylation reaction was set up as follows. FAID14 (1 mg, 100 µL in water) was mixed with 100 µL of 0.1 M Na<sub>2</sub>CO<sub>3</sub> pH 9.6. Biotinyl- $\epsilon$ -amino-caproic acid N-hydroxy succinimide ester (Biotin-X-NHS) (Calbiochem, LaJolla, CA) (2.5 mg) was dissolved in 0.5 mL dimethylsulfoxide. Biotin-X-NHS solution (20 µL) was added to the FAID14 solution and allowed to react at 22°C for 4 hours. Excess biotin and impurities were removed by gel filtration, using a Pharmacia Superose 12 HR10/30 column (Piscataway, NJ). At a flow rate of 0.8 mL/min, the biotinylated FAID14 emerged with a peak at 16.8 min. The fractions making up this peak were pooled and stored at 4°C and used to detect the CC49 idiotype as determined by the CC49  $V_L$  and  $V_H$  CDRs.

#### Isoelectric Focusing (IEF)

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Isoelectric points (pl's) were predicted using a computer program called PROTEIN--TITRATE, available through DNASTAR (Madison, WI). Based on amino acid composition with an input sequence, a MW value is given, in addition to the pl. Since Cys residues contribute to the charge, the count was adjusted to 0 for Cys, since they are all involved in disulfide bonds.

Experimentally, pl's were determined using Isogel agarose IEF plates, pH range 3-10 (FMC Bioproducts, Rockland, ME). A Biorad Bio-phoresis horizontal electrophoresis cell was used to run the IEF, following the directions of both manufacturers. The electrophoresis conditions were: 500 volts (limiting), at 20 mA current and 10 W of constant power. Focusing was complete in 90 min. IEF standards were purchased from Biorad; the kit included phycocyanin, β-lactoglobulin B, bovine carbonic anhydrase, human carbonic anhydrase, equine myoglobin, human hemoglobins A and C, 3 lentil lectins and cytochrome C, with pl values of 4.65, 5.10, 6.00, 6.50, 7.00, 7.10 and 7.50, 7.80, 8.00, and 8.20 and 9.60, respectively. Gels were stained and destained according to the directions provided by FMC.

#### Quantitation of CC49 Antibody Species

All purified CC49 antibodies including the IgG, scFv2 species and the monomeric scFv were quantitated by measuring the absorbence of protein dilutions at 280 mm using matching 1.0 cm pathlength quartz cuvettes (Hellma) and a Perkin-Elmer UV/VIS Spectrophotometer, Model 552A. Molar absorptivities (E<sub>m</sub>) were determined for each antibody by using the following formula:

 $E_{m} = (number Trp) X 5,500 + (number Tyr) X 1,340 + (number (Cys)2) X 150 + (number Phe) X 10$ 

The values are based on information given by D. B. Wetlaufer, Advances in Protein Chemistry, 17, 375-378).

#### High Performance Liquid Chromatography

All high performance liquid chromatography (HPLC) was performed for CC49 scFv2 purification using an LKB HPLC system with titanium or teflon tubing throughout. The system consists of the Model 2150 HPLC pump, model 2152 controller, UV CORD SII model 2238 detection system set at an absorbence of 276 nm and the model 2211 SuperRac fraction collector.

#### **PCR Generation of Subunits**

All polymerase chain reactions (PCR) were performed with a reaction mixture consisting of: 150 picograms (pg) plasmid target (pSCFVUHM); 100 pmoles primers; 1 µL Perkin-Elmer-Cetus (PEC, Norwalk, CT) Ampli-Taq polymerase; 16 µL of 10 mM dNTPs and 10 µL of 10X buffer both supplied in the PEC kit; and sufficient water to bring the volume to total volume to 100 µL. The PCR reactions were carried out essentially as described by the manufacturer. Reactions were done in a PEC 9600 thermocycler with 30 cycles of: denaturation of the DNA at 94°C for 20 to 45 sec, annealing from between 52 to 60°C for 0.5 to 1.5 min., and elongation at 72°C for 0.5 to 2.0 min. Oligonucleotide primers were synthesized on an Applied Biosystems (Foster City, CA) 380A or 391 DNA synthesizer and purified as above.

Ligation reactions using 100 ng of vector DNA and a corresponding 1:1 stoichiometric equivalent of insert DNA were performed using a Stratagene (La Jolla, CA) T4 DNA ligase kit following the manufacturer's directions. Ligation reactions (20 µL total volume) were initially incubated at 18°C and allowed to cool gradually overnight to 4°C.

Transformations

Transformations were performed utilizing 100 µL of Stratagene E. coli AG1 competent cells (Stratagene, La Jolla, CA) according to the directions provided by the manufacturer. DNA from the ligation reactions (1-5 µL) were used. After the transformation step, cells were allowed to recover for 1 hr in Luria broth (LB) at 37°C with continuous mixing and subsequently plated onto either 20 µg/mL chloramphenicol containing (CAM 20) Luria agar for pSCFVUHM, p49LHLH or p49LHHL or 100 µg/mL ampicillin (AMP 100) Luria agar plates

(LB-AMP 100) for clones containing the plasmid pSL301 or subsequent constructions derived from pSL301.

#### Screening of E. coli Clones

Bacterial plasmids were isolated from LB broth culture containing the appropriate drug to maintain selection pressure using Promega (Madison, WI) Magic mini-prep plasmid preparation kits. The kit was used per the manufacturer's specifications.

#### **Plasmid Constructions**

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Two plasmids, designated p49LHLH and p49LHHL, were constructed to produce multivalent single chain antibodies. The host cell containing p49LHLH produced a polypeptide which can be designated by  $V_L$ -L- $V_H$ -L- $V_L$ -L- $V_H$  where  $V_L$  and  $V_H$  are the light and heavy cahin variable regions of CC49 antibody and linker (L) is a 25 amino acid linker having the sequence (SEQ ID NO: 5).

The host cell containing p49LHHL produced a polypeptide which can be designated by  $V_L$ -L- $V_H$ -L- $V_L$  where  $V_L$  and  $V_H$  are the light and heavy chain variable domains of the CC49 antibody and L is a peptide linker having the amino acid sequence indicated above.

The nucleotide sequence (SEQ ID NO: 6) and amino acid sequence (SEQ ID NO: 7) of the CC49  $V_L$ -L- $V_H$ -L- $V_L$ -L- $V_H$  (p49LHLH) are given in Figure 6. The nucleotide sequence (SEQ ID NO: 8) and amino acid sequence (SEQ ID NO: 9) of the CC49  $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_L$  (p49LHHL) are given in Figure 7.

#### Construction of pSL301 HT

The construction of pSL301 HT is illustrated in Figure 8. The Bacillus lichiformist penicillinase P (penP) terminator sequence was removed from the plasmid designated pSCFV UHM by a 45 minute digest with Nhe I and BamH I, excised from a 4.5 percent polyacrylamide gel after electrophoresis, electroeluted, ethanol precipitated and ligated into the same sites in the similarly prepared vector: pSL301 (Invitrogen, San Diego, CA). A procedure for preparing pSCFV UHM is given is U.S. patent application Ser. No. 07/935,695 filed August 21, 1992, the disclosure of which is hereby incorporated by reference. In general, pSCFV UHM contains a nucleotide sequence for a penP promoter; a unique Nco I restriction site; CC49 V<sub>L</sub> region; Hind III restriction site; a 25 amino acid linker; a unique a Xho I restriction site; CC49 V<sub>H</sub> region; Nhe I restriction site; penP terminator; and BamH I restriction site (see, Figure 8). The penP promoter and terminator are described in Mezes, et al. (1983), *J. Biol. Chem.*, 258, 11211-11218 (1983).

An aliquot of the ligation reaction (3 µL) was used to transform competent *E. coli* AG1 cells which were plated on LB-AMP100 agar plates and grown overnight. Potential clones containing the penP terminator insert were screened using a Pharmacia (Gaithersburg, MD) T7

Quickprime <sup>32</sup>P DNA labeling kit in conjunction with the microwave colony lysis procedure outlined in Buluwela et al., *Nucleic Acid Research*, <u>17</u>, 452 (1989). The probe, which was the penP-Nhe I-BamH I terminator fragment itself was prepared and used according to the directions supplied with the Quickprime kit. A clone which was probe positive and which contained the 207 base pair inserts from a BamH I and Nhe I digest (base pairs (bp) 1958 to 2165, Figure 6) was designated pSL301 T and chosen to construct pSL301 HT which would contain the nucleotide sequence for CC49 V<sub>H</sub>. The reason the Nhe I-BamH I penP terminator was placed into pSL301 was to eliminate the Eco47 III restriction endonuclease site present in the polylinker region between its Nhe I and BamH I sites. This was designed to accommodate the subsequent build-up of the V<sub>L</sub> and V<sub>H</sub> domains where the Eco47 III site needed to be unique for the placement of each successive V domain into the construction. As each V domain was added at the Eco47 III-Nhe I sites, the Eco47 III was destroyed in each case to make the next Eco47 III site coming in on the unique insert.

The V<sub>H</sub> sequence was made by PCR with oligos 5' SCP1 and 3'oligo SCP5 using pSCFV UHM as the target for PCR amplification. The DNA sequence for SCP1 (SEQ ID NO: 10) and SCP5 (SEQ ID NO: 11) are as follows:

SCP1: 5'-TAAA CTC GAG GTT CAG TTG CAG CAG -3'

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SCP5: 5'-TAAA GCT AGC ACCA AGC GCT TAG TGA GGA GAC GGT GAC TGA GGT-3'
The underlined portion indicates the endonuclease restriction sites.

The amplified  $V_H$  DNA was purified from a 4 percent PAG, electroeluted ethanol precipitated and dissolved in 20  $\mu$ L water. The  $V_H$  sequence was digested with Xho I and Nhe I restriction enzymes and used as the insert with the pSL301 T vector which had been digested with the same restriction enzymes and subsequently purified. A standard ligation reaction was done and an aliquot (4  $\mu$ L) used to transform competent E. coli AG1 cells. The transformed cells were plated onto LB AMP100 agar plates. Candidate clones were picked from a Nhe I and Xho I digest screen that revealed that the CC49V<sub>H</sub> insert had been obtained.

DNA sequencing was performed to verify the sequence of the CC49V<sub>H</sub> with United States Biochemical (USB) (Cleveland, Ohio) Sequence kit and sequencing primers pSL301SEQB (a 21 bp sequencing primer which annealed in the pSL301 vector 57 bp upstream from the Xho I site) and CC49VHP, revealed clones with the correct CC49V<sub>H</sub> sequence in pSL301HT. This plasmid was used as the starting point in the construction of both pSL301-HLHT and pSL301-HLHT. The sequencing oligos used are shown here.

The nucleotide sequence of pSL301SEQ B (SEQ ID NO: 12) and CC49V $_{\rm H}$  (SEQ ID No: 13) are as follows:

pSL301SEQB: 5'-TCG TCC GAT TAG GCA AGC TTA-3'
CC49VHP: 5'-GAT GAT TTT AAA TAC AAT GAG-3'

#### Example 1 p49LHHL Construction

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Using pSL301 HT (5 µg) as the starting material, it was digested with Eco47 III and Nhe I and the larger vector fragment was purified. A CC49V<sub>H</sub> insert fragment was generated by PCR using SCP6C as the 5' oligo and SCP5 as the 3' oligo. The nucleotide sequence (SEQ ID NO: 14) of SCP6B is as follows:

5'- TAAA TGC GCA GAT GAC GCA AAG AAA GAC GCA GCT AAA AAA GAC GAT GCC AAA AAG GAT GAC GCC AAG AAA GAT CTT GAG GTT CAG TTG CAG CAG TCT-G'

The oligo SCP6B also contains part of the coding region for the linker (bp 8-76 of SEQ ID NO: 14). The portion of the oligo designed to anneal with the CC49VH target in pSCFV UHM is from bp77-90 in SEQ ID NO: 14.

The underlined sequence corresponds to the Fsp I site. The resulting PCR insert was purified, digested with Fsp I and Nhe I and used in a ligation reaction with the pSL301 HT Eco47 III-Nhe I vector (Figure 7). Competent E. coli AG1 cells were used for the transformation of this ligation reaction (3  $\mu$ L) and were plated on LB-AMP100 agar plates. Two clones having the correct size Xho I-Nhe I insert representative of the pSL301 HHT product were sequenced with the oligo SQP1 and a single clone with the correct sequence (nucleotides 1124-1543 of Figure 7) was chosen for further construction. The nucleotide sequence of SQP1 (SEQ ID NO: 16) is as follows:

SQP1: 5'-TG ACT TTA TGT AAG ATG ATG T-3'

The final linker- $V_L$  subunit (bp 1544-1963, Figure 7) was generated using the 5'oligo, SCP7b and the 3' oligo, SCP8a, using pSCFV UHM as the target for the PCR. The nucleotide sequence of SCP7b (SEQ ID NO: 17 is as follows:

SCP7b: 5'-TAAA <u>TGC GCA GAT GAC GCA AAG AAA GAC GCA GCT AAA AAA GAC GAT</u>

GCC AAA AAG GAT GAC GCC AAG AAA GAT CTT GAC ATT GTG ATG TCA CAG TCT

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The underlined nucleotides correspond to an Fsp I site. The nucleotide sequence of SCP8a (SEQ ID NO: 18) is as follows:

SCP8a: 5'-TAAA GCT AGC TTT TTA CTT AAG CAC CAG CTT GGT CCC-3'

The first set of underlined nucleotides correspond to an Nhe I site, while the other corresponds to an Afl II site. Nucleotides 8-76 of SCP70 code for the linker (nucleotides 1544-1612 of Figure 7) while nucleotides 77-99 which anneal to the V<sub>L</sub> correspond to 1613-1635 of Figure 7. The primer SCP8a has a short tail at its 5' end, a Nhe I restriction site, a stop codon, an Afl II restriction site and the last 21 bases of the V<sub>L</sub>. After Fsp I and Nhe I digestion, this resulting 420 bp insert was purified and ligated into the Nhe I and Eco47 III sites of the purified pSL301HHT vector, candidate clones were screened with Nhe I and Xho I, the correct size insert verified and sequenced with 49LFR2(-) and SQP1 to confirm the newly inserted sequence in pSL301HHLT. The nucleotide sequence (SEQ ID NO: 19) is as follows:

#### 49LFR2(-): 5'-CTG CTG GTA CCA GGC CAA G-3'

The plasmid pSL301HHLT was digested with Xho I and Nhe I, purified, and the resulting 1179 bp  $V_H$ -linker- $V_H$ -linker- $V_L$  segment ligated into pSCFV UHM, which had been cut with the same restriction enzymes and the larger vector fragment purified, to form p49LHHL.

The ligation reaction (4 µL aliquot) was used to transform competent E. coli AG1 cells (Stratagene) and plated onto LBCAM20 agar plates. A single clone which had a plasmid with the correct restriction enzyme map was selected to contain p49LHHL. The p49LHHL contains a penP promoter and a nucleotide sequence for the CC49 multivalent single chain antibody scFv2:

 $V_L$ -L- $V_H$ -L- $V_L$  or CC49 scFv2 (LHHL).

Example 2: p49LHLH Construction

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The construction of p49LHLH is schematically represented in Figure 11. A linker- $V_L$  subunit was generated with the 5' oligo SCP7b and the 3'oligo SCP9.

SCP9: 5'-TAA AGC TAG CAC CAA GCG CTT AGT TTC AGC ACC AGC TTG GTC CCA G-3'
The SCP7b oligo (nucleotides 8-76) codes for the linker in Figure 6 (corresponding to nucleotides 1124-1192) and annealed to the pSCFV UHM target for the PCR (nucleotides 77-99) corresponding to nucleotides 1193-1215 of the V<sub>i</sub> in Figure 6.

underlined nucleotides) which are restriction sites needed for making the pSL301HLT ready to accept the next V domain. Nucleotides 18-23 of SCP9 correspond to nucleotides 1532-1537 of Figure 6 (coding for the first 2 amino acids of the linker), while nucleotides 24-46 correspond to nucleotides 1508-1531 of Figure 6 which was also the annealing region for SCP9 in the PCR. The plasmid pSL301 HT was digested with Eco47 III and Nhe I and the larger vector fragment was purified for ligation with the linker-CC49V<sub>L</sub> DNA insert fragment from the PCR which had been treated with Fsp I and Nhe I and purified. The ligation mixture (3 µL) was used to transform *E. coli* AG1 competent cells and one colony having the correct Xho I-Nhe I size fragment was sequenced using the oligo PENPTSEQ2. The nucleotide sequence (SEQ. ID NO. 21) is as follows:

#### 5'-TTG ATC ACC AAG TGA CTT TAT G-3'

The sequencing results indicated that there had been a PCR error and deletion in the resulting pSL301HT clone. A five base deletion, corresponding to nucleotides 1533-1537 as seen in Figure 6 had been obtained and nucleotide 1531 which should have been a T was actually a G, as determined from the DNA sequence data. The resulting sequence was

#### 5'...G AAGC GCT T...etc.

where the underlined sequence fortuitously formed an Eco47 III site. The AGCGCT sequence in Figure 6, would correspond to nucleotides 1530, 1531, 1532, 1538, 1539 and 1540. This error was corrected in the next step, generating pSL301 HLHT, by incorporating the 5 base deletion at the end of oligo SCP6C.

#### SCP6C: 5'-TAAGCGCTGATGATGCTAAGAAGGACGCCGCAAAAAA GGACGACGCAAAAAAAGATGATGCAAAAAAGGATCTGG AGGTTCAGTTGCAGCAGTCTGAC-3'

The underlined sequence in SCP6c corresponds to an Eco47 III site. SCP6C was used as the 5' oligo, with SCP10 as the 3' oligo in a PCR to generate a linker CC49  $V_L$  segment. The nucleotide sequence (SEQ ID NO: 23) is as follows:

SCP10: 5'TTG TGC TAG CTT TTT ATG AGG AGA CGG TGA CTG AGG TT-3'

The underlined sequence in SCP10 corresponds to the Nhe I site found at nucleotides 1958-1963 in Figure 6. The PCR insert was digested this time only with Nhe I and purified. The vector (pSL301 HLT) was digested at the Eco47 III site (that had been formed) and Nhe I and purified. The insert and vector were ligated and an aliquot (3 µL) used to transform competent E. coli AG1 cells. This was plated on LB-AMP100 plates and candidate clones screened with Xho I and Nhe I. Three clones having the correct size DNA were obtained. Two of these clones were sequenced using the oligo 49VLCDR3(+) and SQP1. The nucleotide sequence (DWQ ID NO: 24 of 49VLCDR3(+) is as follows:

49VLCDR3(+):

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5'-CAG CAG TAT TAT AGC TAT-3'

One clone, with the correct sequence was obtained and the sequence from nucleotides 1533 to 1963 in Figure 6 were verified, giving a correct pSL301 HLHL clone.

To generate the final plasmid, p49LHLH for expression in E. coli, pSL301 HLHT (5 μg) was digested with Nhe I and Xho I, and the smaller insert fragment containing the V<sub>H</sub>-L-V<sub>L</sub>-L-V<sub>H</sub> sequence purified. It was ligated with the larger purified vector fragment from a digest of pSCFV UHM (5 μg) with Xho I and Nhe I. An aliquot of the ligation mix (4 μL) was used to transform competent E. coli AG1 cells. The transformation mix was plated on LB-CAM20 plates, and a representative clone for p49 LHLH was selected on the basis of a correct restriction enzyme map (see Figure 10) and biological activity toward TAG-72.

Example 3: Purification of CC49 scFv2 LHLH and LHHL Covalently Linked Dimers

For the purification of the CC49 covalently linked single chain dimers, (scFv2),

E. coli periplasmic fractions were prepared from 1.0 L overnight cultures of both p49LHLH and
p49LHHL. Briefly, the culture was divided into 4 X 250 mL portions and centrifuged at
5,000 rpm for 10 minutes in a Sorvall GS-3 rotor. The pelleted cells were washed and
resuspended in 100 mL each of 10 mM Tris-HCl pH 7.3 containing 30 mM NaCl. The cells were
again pelleted and washed with a total of 100 mL 30 mM Tris-HCl pH 7.3 and pooled into one
tube. To this, 100 mL of 30 mM Tris-HCl pH 7.3 containing 40 percent w/v sucrose and 2.0 mL of
10 mM EDTA pH 7.5 was added. The mixture was rept at room temperature, with occasional
shaking, for 10 minutes. The hypertonic cells were then pelleted as before. In the next step, the
shock, the pellet was quickly suspended in 20 mL ice cold 0.5 mM MgCl<sub>2</sub> and kept on ice for 10
minutes, with occasional shaking. The cells were pelleted as before and the supernatant

containing the *E. coli* periplasmic fraction was clarified further by filtration through a 0.2 µm Nalge (Rochester, NY) filter apparatus and concentrated in Amicon (Danvers, MA) Centriprep 30 and Centricon 30 devices to a volume of less than 1.0 mL.

The concentrated periplasmic shockates from either the p49LHLH or p49LHHL clones were injected onto a Pharmacia (Piscataway, NJ) Superdex 75 HR 10/30 HPLC column that had been equilibrated with PBS. At a flow rate of 0.5 mL/minute, the product of interest, as determined by competition ELISA, had emerged between 21 through 24 minutes. The active fractions were pooled, concentrated as before and dialyzed overnight using a system 500 Microdialyzer Unit (Pierce Chemical) against 20 mM Tris-HCl pH 7.6 with 3-4 changes of buffer and using an 8,000 MW cut-off membrane. The sample was injected on a Pharmacia Mono Q HR 5/5 anion exchange HPLC column. A gradient program using 20 mM Tris-HCl pH 7.6 as buffer A and the same solution plus 0.5 M NaCl as buffer B was employed at a flow rate of 1.5 mL/min. The products of interest in each case, as determined by competition ELISA, emerged from the column between 3 and 4 minutes. Analysis of the fractions at this point on duplicate SDS-PAGE gels, one stained with Coomassie Brilliant Blue R-250 and the other transferred for Western analysis (using biotinylated FAID 14 as the probe antibody) revealed a single band at the calculated molecular weight for the scFv2 (LHLH or LHHL) species at 58,239 daltons. The active fractions were in each case concentrated, dialysed against 50 mM MES pH 5.8 overnight and injected on a Pharmacia Mono S HR 5/5 cation exchange column. The two fractions of interest from this purification step, as determined by SDS-PAGE and ELISA, fractions 5 and 6, eluted just before the start of the gradient, so they had not actually bound to the column. Fractions 5 and 6 were consequently pooled for future purification.

A Mono Q column was again run on the active Mono S fractions but the buffer used was 20 mM Tris-HCl, pH 8.0 and the flow rate was decreased to 0.8 mL/minute. The products emerged without binding, but the impurity left over from the Mono S was slightly more held up, so that separation did occur between 5 and 6 minutes. After this run, the products were homogeneous and were saved for further characterization.

#### Isoelectric Focusing

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The isoelectric points (pl) of the constructs was predicted using the DNASTAR (Madison, WI) computer program Protein-titrate. Based on amino acid composition, a MW and pl value was calculated.

Experimentally, pls were determined using FMC Bioproducts (Rockland, ME) Isogel IEF plates, pH range 3-10. A Biorad (Richmond, CA) electrophoresis unit was used to run the IEF, following the directions of both manufacturers. The electrophoresis conditions were as follows: 500 V (limiting) at 20 mA and at 10 W of constant power. Focusing was complete in 90 minutes. Biorad IEF standards included phycocyanin, beta lactoglobulin B, bovine carbonic anhydrase, human carbonic anhydrase, equine myoglobulin, human hemoglobins A and C, 3 lentil lectin, and cytochrome C with pI value of 4.65, 5.10, 6.00, 6,50, 7.00, 7.50, 7.8, 8.00, 8.20

and 9.6, respectively. Gels were stained and destained according to directions provided by FMC. The DNASTAR program predicted values of 8.1 for the pl for both scFv2 species. A single, homogeneous band for the pure products was observed on the gel at pl values for both at 6.9.

Purified CC49 antibodies such as the IgG, scFv2 (LHLH and LHHL) were quantitated by measuring the absorbence spectrophotometrically at 280 nm. Molar absorbtivity values,  $\epsilon_{\rm M}$ , were determined for each using the formula cited above by Wetlaufer.

Based on the amino acid composition, the E<sup>0.1%</sup> (280 nanometers) values for CC49 IgG, CC49 scFv2 LHLH, CC49 scFv2 LHHL and CC49 scFv were 1.49, 1.65, 1.65 and 1.71, respectively.

#### 10 Example 4

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Relative activities of the CC49 scFv2 species LHLH and LHHL, were compared with the IgG and a monomer scFv form with a FLAG peptide at the COOH terminus.

Percent competition was determined from the ELISA data by the following equation:

Zero competition - sample reading (OD405-450 nm)  $\times$  100 zero competition - 100 percent competition

The "zero competition" value was determined by mixing (1:1) one percent BSA with the biotinylated CC49 (3 X 10-14 moles) while the 100 percent competition value was based on a 5 µg/mL sample of CC49 IgG mixed with the biotinylated CC49 IgG. The data are presented in Figure 11. Absorbence values for the samples were measured at 405 nm - 450 nm. The average of triplicate readings was used. Initially samples (25 µL) were applied to the TAG-72 coated microliter plates at 1.0 X 10-10 moles of binding sites/mL. Biotinylated CC49 (4 µg/µL diluted 1:20,000 - used 25 µL) diluted the samples by a factor of 2. Serial dilutions (1:2) were performed. Both forms of the scFv2 are approximately equivalent to the IgG (see Figure 11). In a separate experiment, a CC49 scFv monomer was compared to a Fab fragment, both of which are monovalent and these were also shown to be equivalent in their binding affinity for TAG-72. These results indicate that both forms of the covalently linked dimers have 2 fully functional antigen binding sites. This is the same increase in avidity as observed with the whole IgG, relative to a monomeric species.

These data also indicate that the scFv2 molecules, like their CC49 IgG parent are candidates for immunotherapeutic applications, but with the benefit of increased capillary permeability and more rapid biodistribution pharmacokinetics. The advantage should allow multiple injections of compounds of the present invention and give higher tumor: tissue ratios in immunotherapeutic treatment regimens for cancer treatment, relative to the existing IgG molecules.

Other embodiments of the invention will be apparent to those skilled in the art from a consideration of this specification or practice of the invention disclosed herein. It is

intended that the specification and examples be considered as exemplary only, with the true-scope and spirit of the invention being indicated by the following claims.

WO 94/13806

PCT/US93/12039

1. A mutivalent single chain antibody which comprises two or more single chain antibody fragments each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:

- (a) a first polypeptide comprising a light chain variable domain;
- (b) a second polypeptide comprising a heavy chain variable domain; and
- (c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.
- 2. The multivalent single chain antibody of Claim 1 wherein the first peptide linker has the amino and sequence

Leu Ser Ala Asp Asp Ala Lys Lys Asp Ala Lys Lys Asp Ala Lys Lys Asp Ala Lys Lys Asp Leu.

- 3. The multivalent single chain antibody of Claim 1 wherein the light chain variable region has an amino acid sequence substantially the same as that of Figure 3 and the heavy chain variable region has an amino acid sequence substantially the same as that of Figure 5.
- 4. The multivalent single chain antibody of Claim 1 wherein the first and second peptide linkers have an amino acid sequence which is substantially the same.
- 5. A DNA sequence which codes for a mutivalent single chain antibody, the multivalent single antibody comprising two or more single chain antibody fragments, each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:
  - (a) a first polypeptide comprising a light chain variable domain;
  - (b) a second polypeptide comprising a heavy chain variable domain; and
  - (c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.
- 6. The DNA sequence of Claim 5 wherein the sequence coding for the first polypeptide is substantially the same as that of Figure 2 and the sequence coding for the second polypeptide is substantially the same as that of Figure 3.

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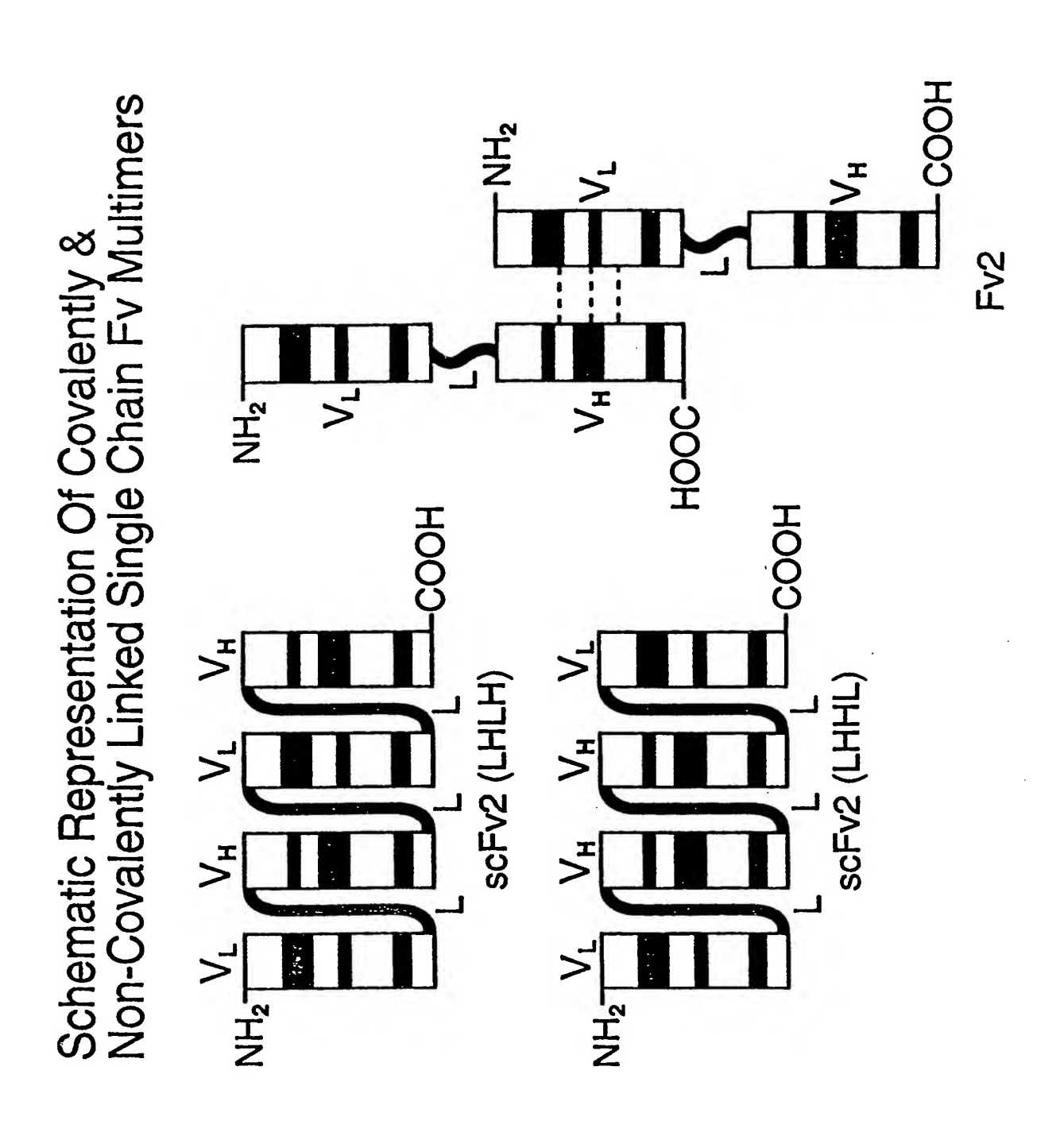
25

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FIGURE 1



### FIGURE 2

GAC ATT GTG ATG TCA CAG TCT CCA TCC TCC CTA CCT GTG TCA GTT GGC GAG AAG GTT ACT TTG AGC TGC AAG TCC AGT CAG AGC CTT TTA TAT AGT GGT AAT CAA AAG AAC TAC TTG GCC TGG TAC CAG CAG AAA CCA GGG CAG TCT CCT AAA CTG CTG ATT TAC TGG GCA TCC GCT AGG GAA TCT GGG GTC CCT GAT CGC TTC ACA GGC AGT GGA TCT GGG ACA GAT TTC ACT CTC TCC ATC AGC AGT GTG AAG ACC TAT TAT TAT AGC TAT CCC CTC ACG CAG TAT TAT AAGC TAT CCC CTC ACG TTC ACG GTG CTG AAG

### FIGURE 3

Asp Ile Val Met Ser Gln Ser Pro Ser Ser Leu Pro Val Ser Val Gly Glu Lys Val Thr Leu Ser Cys Lys Ser Ser Gln Ser Leu Leu Tyr Ser Gly Asn Gln Lys Asn Tyr Leu Ala Trp Tyr Gln Gln Lys Pro Gly Gln Ser Pro Lys Leu Leu Ile Tyr Trp Ala Ser Ala Arg Glu Ser Gly Val Pro Asp Arg Phe Thr Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Ser Ile Ser Ser Val Lys Thr Glu Asp Leu Ala Val Tyr Tyr Cys Gln Gln Tyr Tyr Ser Tyr Pro Leu Thr Phe Gly Ala Gly Thr Lys Leu Val Leu Lys

### FIGURE 4

GAG GTT CAG TTG CAG CAG TCT GAC GCT GAG TTG GTG AAA CCT
GGG GCT TCA GTG AAG ATT TCC TGC AAG GCT TCT GGC TAC ACC
TTC ACT GAC CAT GCA ATT CAC TGG GTG AAA CAG AAC CCT GAA
CAG GGC CTG GAA TGG ATT GGA TAT TTT TCT CCC GGA AAT GAT
GAT TTT AAA TAC AAT GAG AGG TTC AAG GGC AAG GCC ACA CTG
ACT GCA GAC AAA TCC TCC AGC ACT GCC TAC GTG CAG CTC AAC
AGC CTG ACA TCT GAG GAT TCT GCA GTG TAT TTC TGT ACA AGA
TCC CTG AAT ATG GCC TAC TGG GGT CAA GGA ACC TCA GTC ACC
GTC TCC TCA

### FIGURE 5

Glu Val Gln Leu Gln Gln Ser Asp Ala Glu Leu Val Lys Pro Gly Ala Ser Val Lys Ile Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Asp His Ala Ile His Trp Val Lys Gln Asn Pro Glu Gln Gly Leu Glu Trp Ile Gly Tyr Phe Ser Pro Gly Asn Asp Asp Phe Lys Tyr Asn Glu Arg Phe Lys Gly Lys Ala Thr Leu Thr Ala Asp Lys Ser Ser Ser Thr Ala Tyr Val Gln Leu Asn Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Phe Cys Thr Arg Ser Leu Asn Met Ala Tyr Trp Gly Gln Gly Thr Ser Val Thr Val Ser Ser

## FIGURE

DNA AND AMINO ACID SEQUENCE OF CC49 VL-L-VH-L-VL-L-VH

46	ħ6	142	190	238	286	334	382	430	478
TCC	AAA	ATT	TIA	AGA	ACG	Leu TTA	Pro	Lys	40 Ala GCC
၁၁၁	GAA	TAC	CAA	GAA	GAG	Leu	Ser	Cys	Leu TTG
CIT	TCC	GAC	TAC	GTG	AGG	Leu TTG	Gln CAG	Ser	Tyr
TCA	CCT	AAA	ACG	ATA	<b>9</b> 90	Gly GGA	Ser TCA	Leu TTG	Asn
CCA	TTT	TTC	CGT	999 999	AAA TTT	Ala GCT	Met	20 Thr ACT	Lys
ATT	TCA	ACT	GAT	GTA	TTC	<b>HO</b>	Val	Val	Gln
Eco TGA	TCA	AAT	AAA	ACT	ATA TAT	Ala GCA	Ile	Lys	Asn
CGA	AGG	TAT	CTG	AAC	AT CAA ENPR2-	Thr	VL Asp GAC	Glu GAG	Gly GGT
Cla I CAT CGA	ACG	ATA	550	R1- CAT	AAT PENF	Pro	Ala GCC	Gly	Ser
TAT	GAA ACG AGG	TAC	AGT	PENP	ATC	Leu TTG	Met ATG	Val	Tyr
GCT	GTG	TCT	TIG	TCA	ACG	Leu	Nco Ala GCC	Ser	30 Leu TTA
TTG ACA	ອວວ	AAA	TGT	TGT	GTT	Tyr	Pro	Val	Leu
TTG	TCC CCG	TTT	TGA TGT	GAT	CTG	Lys AAA	Gln	Pro	Ser
TGT	TIG	GCA	ATT	CGT	CAT	-22 Met ATG	Ala GCC	Leu	Gln
TCA	CAT	GTT	AAG ATT	TTT	CTT	TIG	Ala GCT	Ser TCC	Ser
51-0	GTT	ACG	TGT	TTG	GTG	ATT	Leu	Ser	Ser

# 'IGURE 6 (Cont. (2))

526	574	622	670	718	166
Trp	G1y GGA	Asp	Phe	120 Lys AAA	Lys
Tyr	Ser	Glu GAA	Thr	Ala GCG	Lys
Ile	70 G1y GGC	Thr	Cic	Asp GAT	Ala GCT
Leu	Thr	Lys	970 000	Asp	Asp
Leu	Phe	Val GTG	100 Tyr TAT	Ala GCG	Asp
Lys	Arg	Ser	Ser	Ser	Lys
50 Pro CCT	Asp Gat	Ser	Tyr	Leu	130 Lys AAG
Ser	Pro	Ile	Tyr	Hind Lys AAG	Ala GCT
Gln	Val	Ser TCC	Gln	Leu	Asp
Gly	GLY	Leu	Gln	Val	Asp
Pro	Ser	Thr	Cys	110 Leu CTG	Lys
Lys AAA	Glu	Phe	Tyr	Lys	Lys
Gln CAG	60 Arg Agg	Asp GAT	Tyr	Thr	Ala GCG
Gln CAG	Ala GCT	Thr	Val	GIY	Ala GCT
Tyr	Ser	G1y GGG	90 Ala GCA	Ala GCT	Asp
Trp TGG	Ala GCA	Ser	Leu CTG	Gly	Lys

# FIGURE 6 (Cont. (3))

814	862	910	958	1006	1054
Pro	Thr	Glu	GAGGAG	Thr	Tyr
Lys	Phe	Leu	AAT Asn AAT	Ser	Val
150 Val	Thr	Gly	200 TAC Tyr	Ser	230 Ala GCA
Leu G	Tyr	Gln	AAA Lys Aaa	Ser	Ser
Glu 1G TT	Gly	180 Glu GAA	TTT Phe TTT	Lys	Asp Gat
Ala T GA	Ser	Pro	GAT ASP GAT	Asp GAC	Glu
Asp C GC	Ala GCT	Asn	GAT Asp GAT	210 Ala GCA	Ser
Ser CI GA	Lys	Gln	Asn AAT	Thr	Thr
Gln AG T(	160 Cys TGC	Lys	CC#9V Gly GGA	Leu CTG	Leu
Gln AG C	Ser	Val GTG	S C C C C C C C C C C C C C C C C C C C	Thr	Ser
Leu TG C	Ile ATT	Trd	190 Ser TCT	Ala GCC	Asn
Gln AG TJ	Lys	His	Phe TTT	Lys	Leu
140 Val TT C/	Val	Ile ATT	Tyr	Gly	220 Gln CAG
1 G1 G G	Ser	Ala GCA	Gly	Lys	Val
Xbo Lec GA	Ala GCT	170 His CAT	Ile	Phe TTC	Tyr
Asp GAC CT	GIY	ASP	Trp	Arg	Ala

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# 'IGURE 6 (Cont. (4))

1102	1150	1198	1246	1294	1342
Ser	Ala GCT	280 11e ATT	Lys	Asn	Pro
Thr	Ala GCA	VL Asp GAC	Glu	Gly	Ser
G G G G G G	Asp	Leu	GIY	310 Ser AGT	Gln CAG
CAA Gln CAA	Lys AAA	Asp	Val	Tyr	Gly
667 667 667	260 Lys AAG	Lys AAA	Ser	Leu TTA	Pro
766 777 766	Ala GCA	Lys	Val	Leu	Lys AAA
TAC TAC TAC	Asp	Ala	290 Pro CC1	Ser	Gln CAG
GCC GCC GCC	Asp Gat	ASP	Leu	Gln	Gln CAG
240 ATG Met ATG	Ala GCA	Asp	Ser	Ser	320 Tyr TAC
AAT Asn AAT	Ser AGC	Lys	Ser	Ser	14 160
Le G CTG	Leu	270 Lys AAA	Pro	Lys	Ala GCC
H49J Ser TCC	Ser	Ala	Ser	CY TGC	Leu
Arg AGA	Ser	Asp	Gln	300 Ser AGC	Tyr
Thr	Val	Asp	Ser	Leu TTG	Asn
Cys	250 Thr ACC	Lys	Met	Thr	Lys
Phe	Val	Lys	Val	Val	Gln

# 'IGURE 6 (Cont. (5))

1390	1438	1486	1534	1582	1630	1678
Asp	360 Ser AGC	TAT Tyr TAT	Leu	Lys	Asp	440 Ala GCT
Pro	I10 ATC	TAT Tyr TAT	Lys	Ala	Ser	Lys
Val	Ser	CAG Gln CAG	390 Leu CTG	Asp	Gln	Cys TGC
Gly	Leu	CAG Gln CAG	Val	Asp	Gln	Ser
340 Ser TCT	Thr	Cys TGT	Leu	Lys	420 Leu TTG	Ile
Glu GAA	Phe	VLCDR Tyr TAC	Lys	Lys	Gln CAG	Lys
Arg	Asp	370 49V Tyr TAT	Thr	Ala	Val	Val GTĞ
Ala GCT	Thr	Val	G1y GGG	Ala	VH Glu GAG	Ser
Ser	Gly	Ala GCA	Ala GCT	400 Asp GAC	Leu	Ala GCT
Ala GCA	Ser	Leu CTG	Gly	Lys	Asp Gat	Gly
Trd TGG	350 Gly GGA	Asp	Phe	Lys	Lys AAG	430 Pro CCT
Tyr	Ser	Glu	Thr	Ala GCT	Lys	Lys
Ile	GLY	Thr	380 Leu CTC	Asp	Ala GCA	Val
Leu	Thr	Lys	Pro	Asp GAT	Asp	Leu TTG
330 Leu CTG	Phe TTC	Val	TAT Tyr TAT	17 II Ala GCT	410 Asp Gat	Glu GAG
Lys	Arg	Ser	AGC Ser AGC	Eco4 Ser AGC	Lys	Ala GCT

# FIGURE 6 (Cont. (6))

1726	1774	1822	1870	1918	1966
Asn	Asp GAT	Ala GCA	Ser	520 Tyr TAC	GAT
Gln	Asn AAT	Thr	Thr	Ala GCC	I AGC
Lys	470 Gly GGA	CTe	Leu	Met	Nhe
Val GTG	Pro CCC	Thr	Ser	Asn	AAA
Trd TGG	Ser	Ala GCC	500 Asn AAC	Leu CTG	*** TAA
His	Phe TTT	Lys	Lec	Ser	Ser
450 Ile ATT	Tyr	Gly	Gln	Arg	530 767 767
Ala GCA	G1y GGA	Lys	Val GTG	Thr	Val
His	Ile	480 Phe TTC	Tyr	Cys	Thr
Asp	Trd	Arg	Ala GCC	Phe TIC	Val
Thr	Glu GAA	Glu	Thr	510 Tyr TAT	Ser
Phe TTC	Leu	Asn	Ser	Val	Thr
Thr	460 Gly GGC	Tyr	Ser	Ala GCA	Gly GGA
Tyr	Gln	Lys	Ser	Ser	Gln
Gly	Glu GAA	Phe TTT	490 Lys AAA	Asp Gat	Gly
Ser	Pro	Asp	Asp	Glu GAG	Hr TGG

### "IGURE 6 (Cont. (7))

GAA	TCC	GIC	GTC AAA	AC/	TCA	TCT	TAC	ATA	AAG	TCA	CTT	GGT	GAT CAA GCT	CAA	GCT	2014
		S(	2P 1-	T.	AGI	FAGT AGA AT PENPTSEQ2-	A I G	TAT	TIC	AGT	GAA	GAA CCA CTA	CTA	GTT		
CAT	ATC		AIT GIC	990	CGG CAA TGG	TGG	TGT	999	CIT	TIT	TIG	TTT	TCT	ATC	TTT	2062
AAA	GAT	CAT	GTG		AAA	AAG AAA AAC GGG AAA	<b>ව</b> වව	AAA	ATC	GGT	CTG	990	CGG GAA AGG ACC	AGG	ACC	2110
999	TTT	TTG	TCG		TCA	TAG	ອວອ	AAT	999	TTG	GAT	GAT TGT GAC AAA ATT	GAC	AAA	ATT	2158
BamH	I Ha	Č														2165

### FIGURE /

DNA AND AMINO ACID SEQUENCE OF CC49 VL-L-VH-L-VL

97	<b>π6</b>	142	190	238	286	334	382	430
rcc	AAA	ATT	TTA	AGA	ACG	ne TA	ro CA	8 G
۲						HE	A O	LY
၁	GAA	TAC	CAA	GAA GAA	GAG	Leu TTA	Ser	Cys
TCA CIT	TCC	GAC	TAC	GTG GTG	AGG	Leu TIG	Gln	Ser
	CCT	AAA	ACG	ATA ATA	ဗဗ္ဗာ ဗဗ္ဗာ	Gly	Ser	Leu
CCA	TTT	TTC	CGT	999 999	AAA TTT	Ala GCT	Met	20 Thr ACT
ATT	TCA	ACT	GAT	GTA	TTC	Ala	Val GTG	Val
EcoR TGA A	TCA	AAT	AAA	ACT	ATA	Ala GCA	Ile	Lys AAG
CGA	AGG	TAT	CTG	AAC	CAA	Thr	VL Asp GAC	Glu
CAT	ACG	TAC ATA TAT	550	CAT	AAT CAA PENPR2-	Pro	Ala GCC	G G G G G G
TAT	GAA ACG AGG	TAC	AGT	PENPR1- AGC CAT	ATC	Leu TTG	Met ATG	Val
TTG ACA GCT TAT	GTG	TCT	TTG	TCA	ACG	Leu	Nco Ala l GCC	Ser
ACA	TCC CCG	TTT AAA	TGT	TGT	GTT	Tyr	Pro	Val
TTG	TCC	TTT	TGA TGT	GAT TGT	CTG	Lys AAA	Gln	Pro
CA TGT	TTG	GCA		CGT	CAT	-22 Met ATG	Ala GCC	Leu
TCA	CAT TTG	GTT	AAG ATT	TTT CGT	CTT CAT	TTG	Ala GCT	10 Ser TCC
5,-0	GTT	ACG	TGT	TIG	GTG	ATT	Leu	Ser

### FIGURE 7 (Cont. (2))

478	526	574	622	670	718	766
40 Ala GCC	Trp TGG	G1y GGA	Asp GAC	<b>Phe</b> TTC	120 Lys AAA	Lys AAG
Leu	Tyr	Ser AGT	Glu GAA	Thr	Ala GCG	Lys
Tyr	Ile	70 617 660	Thr	Leu	Asp	Ala GCT
Asn	Leu	Thr	Lys	Pro	Asp GAC	Asp GAT
Lys	Leu	Phe TTC	Val	100 Tyr TAT	Ala GCG	Asp GAC
Gln	Lys	Arg	Ser	Ser	Ser	Lys
	Pro	Asp GAT	Ser AGC	Tyr	Leu	130 Lys AAG SEQ
Gly	Ser	Pro	Ile	Tyr	Hind Lys AAG	Ala GCT 'L(-)
Ser	Gln	Val	80 Ser TCC	Gln	Leu	Asp GAC TMNV
<b>Tyr</b> TAT	Gly	Gly	Leu	Gln	Val GTG	Asp GAT CTA
30 Leu TTA	Pro	Ser	Thr	Cys	110 Leu CTG	Lys AAG TTC
Leu	Lys AAA	Glu GAA	Phe TTC	Tyr	Lys	Lys AAG TTC
Ser AGC	Gln CAG	60 Are Agg	Asp GAT	Tyr	Thr	Ala GCG CGC
Gln	Gln CAG	Ala GCT	Thr	Val	GIY	Ala GCT CGA
Ser	Tyr	Ser	G1y GGG	90 Ala GCA	Ala GCT	Asp GAT CTA
Ser	Trp TGG	Ala GCA	Ser	Leu	Gly GGT	Lys AAG TTC

### FIGURE 7 (Cont. (3))

814	862	910	958	1006	1054	1102
Pro	Thr	Glu	200 GAG Glu GAG	Thr	Tyr	Ser
Lys AAA	Phe TTC	Leu	AAT Asn AAT	Ser	Val GTG	Thr
150 Val GTG	Thr	Gly	TAC Tyr TAC	Ser	230 Ala GCA	G Gly GGA
Leu TTG	Tyr	Gln	AAA Lys Aaa	Ser	Ser	CAA Gln CAA
Glu GAG	GIY	180 Glu GAA	TTT Phe TTT	Lys	Asp Gat	GGT Gly GGT
Ala GCT	Ser	Pro	GAT Asp GAT	Asp	Glu GAG	166 170 166
Asp	Ala GCT	Asn	GAT Asp GAT	210 Ala GCA	Ser	TAC Tyr TAC
Ser	Lys	Gln	Asn AAT	Thr	Thr	GCC Ala GCC
Gln	160 Cys TGC	Lys	C49V G1y GGA	Leu CTG	Leu	240 ATG Met ATG
Gln CAG	Ser	Val	Pro CCC	Thr	Ser	AAT Asn AAT
Leu	Ile Att	ir iga	190 Ser TCT	Ala GCC	Asn	J- G Leu CTG
Gln	Lys	His	Phe TTT	Lys	Leu	H49 Ser TCC
140 Val GTT	Val	Ile Att	Tyr	GIY	220 Gln CAG	Arg Aga
IGIO	Ser	Ala GCA	Gly	Lys	Val	Thr
Xho Leu CTC	Ala GCT	170 His CAT	Ile	Phe	Tyr	Cys
Asp	<b>G1y</b> GGG	Asp	Trp	Arg	Ala GCC	Phe

## FIGURE 7 (Cont. (4))

1150	1198	1246	1294	1342	1390	1438	1486
Ala GCT	280 Val GTT	Val	Ile ATT	Tyr	GLY	360 Gln CAG	Arg
la CA	VH G1u GAG	Ser	Ala GCA	GIY	Lys	Val	Thr
ASP A GAC G	Leu	Ala	310 His CAT	ILO	Phe TTC	Tyr	Cys
Lys A	Asp GAT	GLY	Asp	Trp	Arg	Ala GCC	Phe TTC
260 Lys I AAG A	Lys	Pro	Thr	Glu	340 G1u GAG	Thr	Tyr
la CA	Lys	Lys	Phe	Cre	Asn	Ser	Val
A G	Ala GCC	290 Val GTG	Thr	GGC	Tyr	Ser	370 Ala GCA
Sp A	Asp	Leu TTG	Tyr	Gln	Lys	Ser	Ser
lla A SCA G	Asp Gat	Glu	Gly	320 Glu GAA	Phe TTT	Lys	Asp Gat
ser A	Lys	Ala GCT	Ser	Proceed	Asp Gat	Asp	Glu
eu S	270 Lys AAA	Asp	Ala GCT	Asn	Asp GAT	350 Ala GCA	Ser
Ser CA C	Ala GCC	Ser	Lys	Gln	Asn	Thr	Thr
Ser S	Asp GAT	Gln	300 Cys 1GC	Lys	Gly	Leu	Leu CTG
ral S	Asp	Gln	Ser	Val	Pro	Thr	Ser
ibr d	Lys AAA	Leu	Ile	Trp	330 Ser TCT	Ala GCC	Asn
Val T GTC A	Lys	Gln	Lys	His	Phe TTT	Lys	Leu

### 'IGURE 7 (Cont. (5))

1534	1582	1630	1678	1726	1774	1822
Ser	Asp	Gln	440 Ser Agc	Tyr TAC	Ile	Gly
Val	Asp	Ser	Leu TIG	Asn AAC (2(-)	Leu	Thr
390 Thr ACC	Lys	Met	Thr	Lys AAG 49LFR	470 Leu CTG	Phe TTC
Val	Lys	Val	Val	Gln	Lys	Arg
Ser	Ala GCT	420 11e ATT	Lys	Asn	Pro	Asp Gat
Thr	Ala GCA	VL Asp GAC	Glu	Gly	Ser	Pro
Gly	Asp	Leu	GLY	450 Ser AGT	Gln	Val
Glu	Lys	Asp	Val	Tyr	Gly	Gly
Gly	400 Lys AAG	Lys	Ser	Leu TTA	Pro	480 Ser TCT
Tr TGG	Ala GCA	Lys	Val	Leu	Lys	Glu GAA
Tyr	Asp	Ala GCC	430 Pro	Ser	Glucato	Arg
Ala	Asp Gat	ASP	Leu	Gln	Glucato	Ala GCT
380 Met ATG	Ala GCA	Asp	Ser	Ser AGT	460 Tyr TAC ATG	Ser
Asn	Ser	Lys	Ser	Ser	Trp TGG ACC	Ala GCA
Leu	Leu	410 Lys AAA	Pro	Lys	Ala GCC CGG	Trp
Ser	Ser	Ala GCC	Ser	Cys	Leu TTG AAC	Tyr

# FIGURE 7 (Cont. (6))

1870	1918	1966	2014		2062	2110	2158	2165
Thr	520 Leu CTC	GAT	GCT		TTT	ACC	ATT	
Lys	Pro	IAGC	CAA	GTT	ATC	AGG	AAA	
Val GTG	Tyr TAT	Nhe	GAT	CTA	TCT	GAA	GAC	٠
Ser	Ser AGC	AAA	GGT	CCA	TIT	990	TGT	
500 Ser AGC	Tyr	*** TAA	CTI	GAA	TIG	CTG	GAT	
Ile	Tyr	II Lys AAG	TCA	AGI	TTT	GGT	TIG	
Ser	Gln CAG	Afil Cer CTT	AAG	110	CTT	ATC	999	
Leu	Gln	Val	_ <b>[</b> → •	TAT	<del>ຽ</del>	AAA	AAT	
Thr	Cys TGT	Leu	TAC	ATG 2- G	TGT	999	909	
<b>Phe</b> TTC	Tyr	Lys	TCT	AGA SEQ	TGG	AAC	TAG	
Asp GAT	510 Tyr TAT	Thr	TCA	PENPT	CAA	AAA	TCA	
Thr	Val	G1y GGG	ACA	GI	550	AAG	AAA	
GIY	Ala GCA	Ala GCT	AA	2P 1-	GIC	GTG	TCG	•
Ser TCT	Leu	Gly	GTC	S	ATT	CAT	TTG	C-3
490 Gly GGA	Asp	Phe	TCC		ATC	GAT	TTT	nH I ATC
Ser	Glu GAA	Thr	GAA		CAT	AAA	555	BamH CGG A

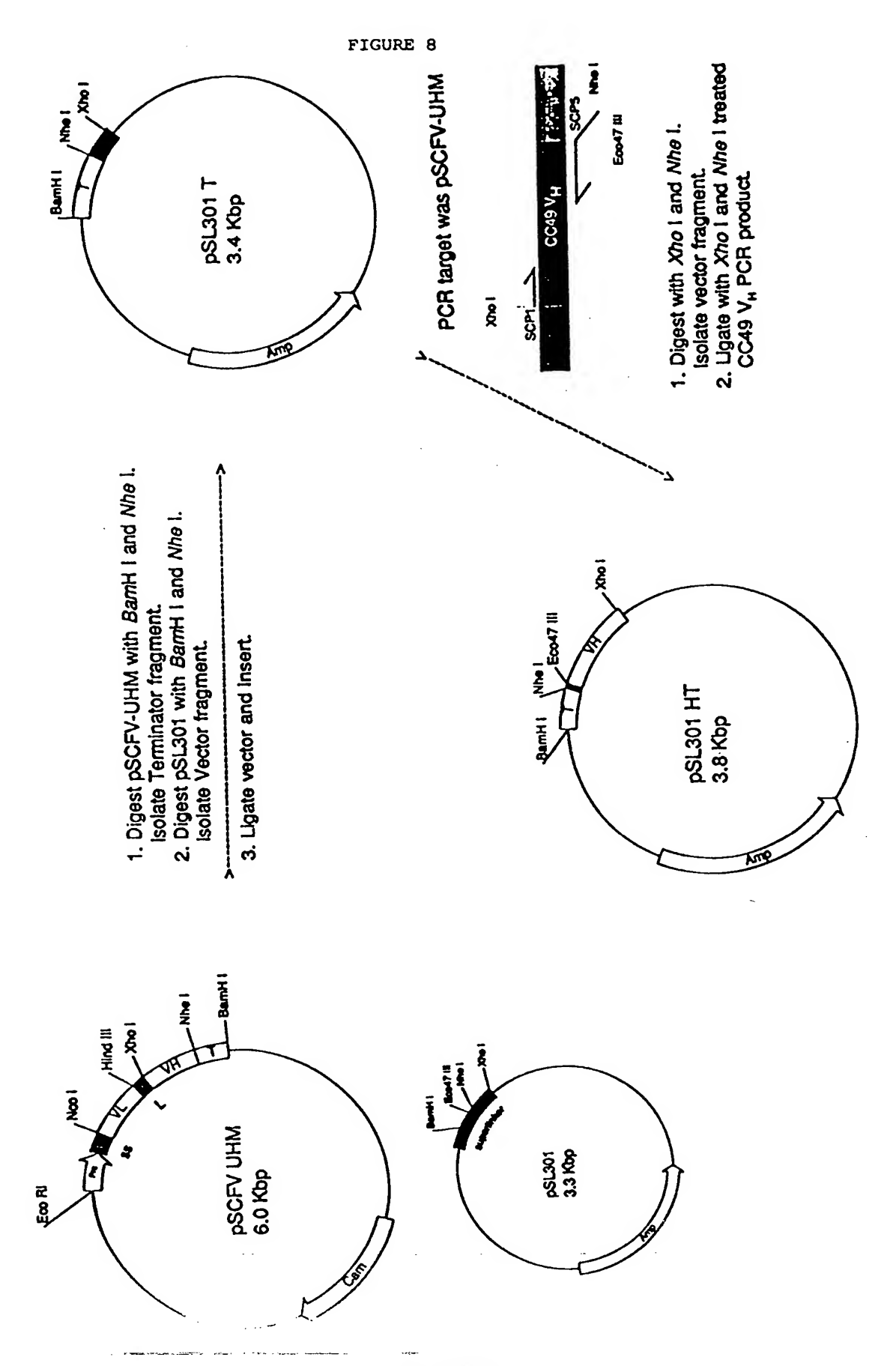
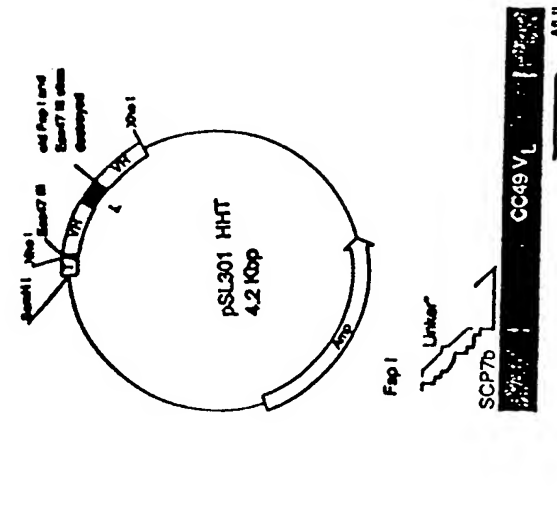


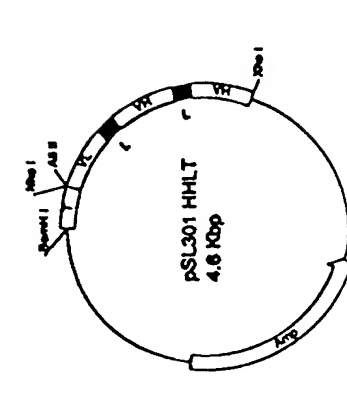
FIGURE 9



SCP84

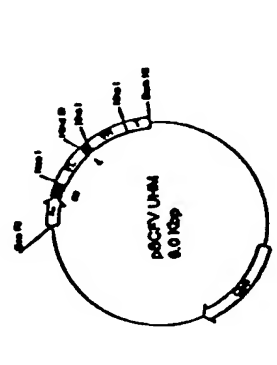
PCR target was pOCPV-URA

Ligate with Fsp I and Nhe I treated CC49 V, PCR product. 1. Digest with Eco47 III and Nhe I. Isolate vector fragment.



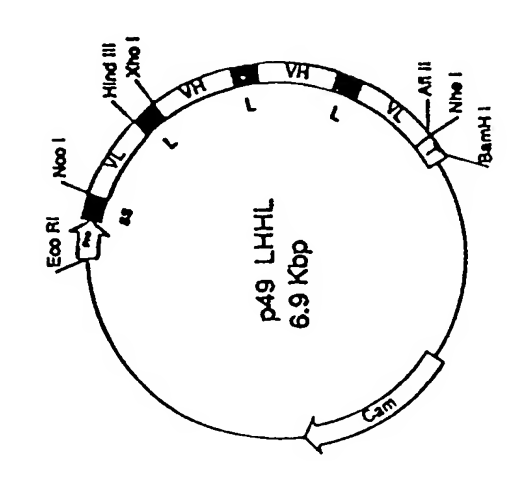
CC49 VH Santos [PASSAS / NATO - <u>\*</u> PCR target was pSCFV-UHM E0047 III

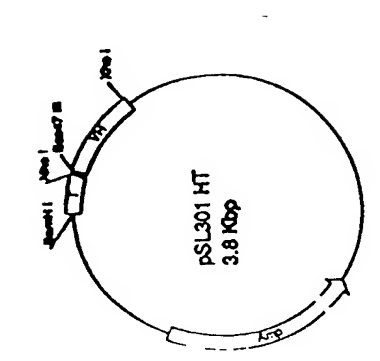
1. Digest with Eco47 III and Nhe I. Isolate vector fragment. and Nhe I treated 2. Ligate with Fsp I and I CC49 V<sub>H</sub> PCR product



1. Digest pSL301 HHLT with Nhe I and Xho I. Isolate HHL fragment.

Isolate vector fragment and ligate with insert. 2. Digest pSCFV-UHM with Xho I and Nhe I





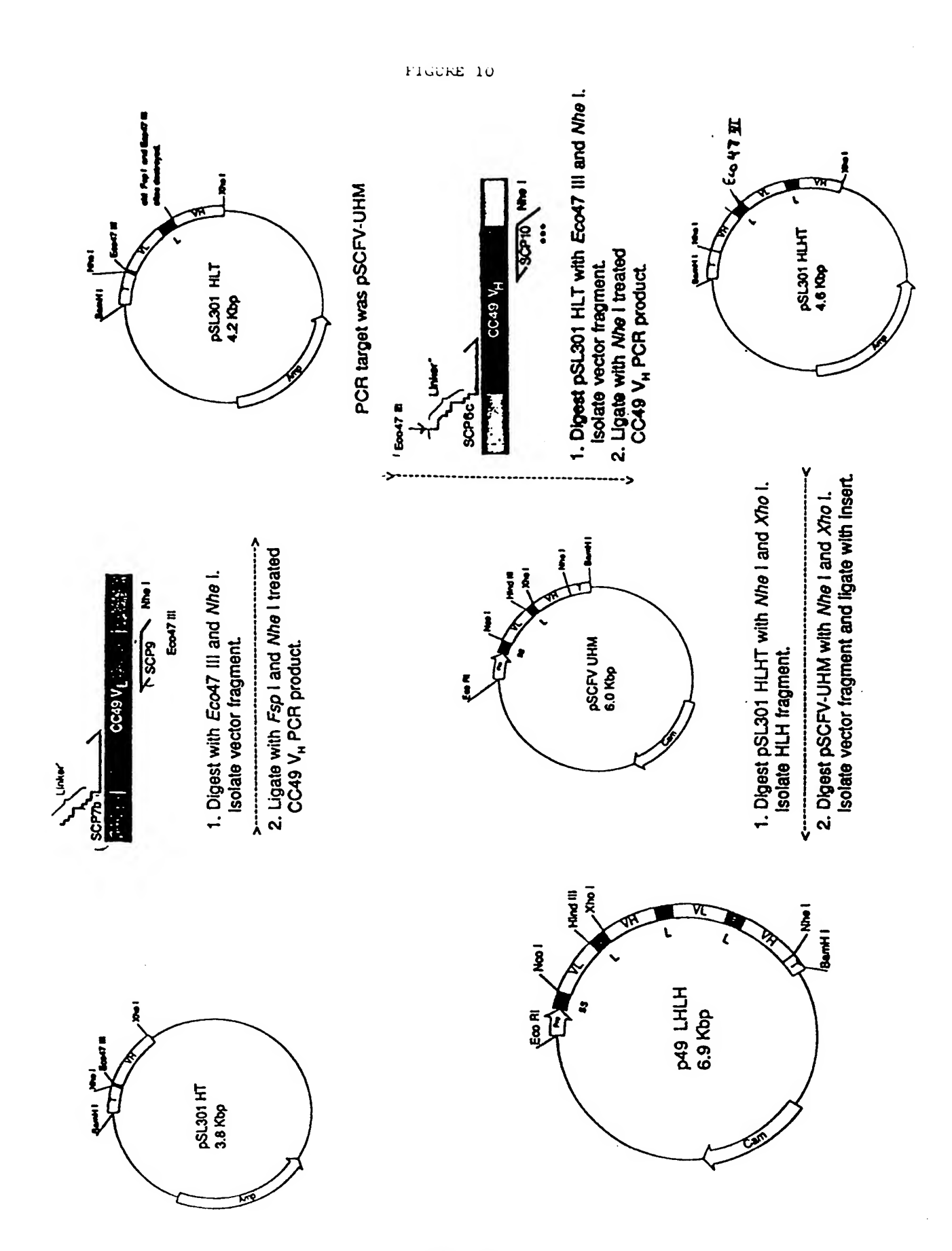
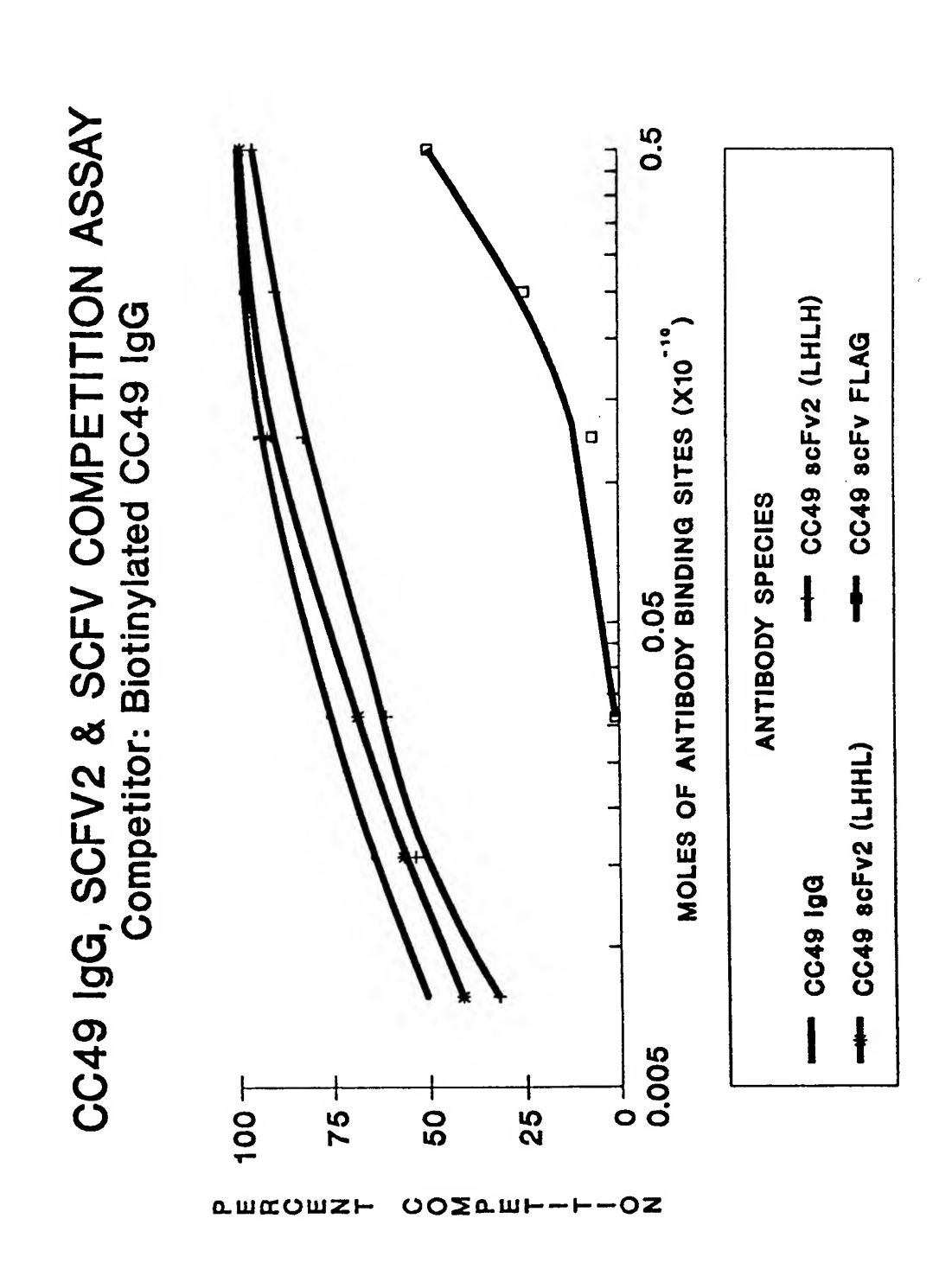


FIGURE 11



### INTERNATIONAL SEARCH REPORT

al Application No Intern

PCT/US 93/12039 A. CLASSIFICATION OF SUBJECT MATTER IPC 5 C12N15/13 C07K15/28 A61K39/395 C12N15/62 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C12N C07K IPC 5 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Category ° Citation of document, with indication, where appropriate, of the relevant passages WO, A, 91 19739 (CELLTECH LIMITED) 26 1,5 X December 1991 2-4,6see example 1 3,6 CANCER RESEARCH vol. 52, no. 12 , 15 June 1992 , PHILADELPHIA, PA, USA pages 3402 - 3408 T.YOKATA ET AL. 'Rapid tumour penetration of a single-chain Fv and comparison with other immunoglobulin forms' see page 3403, column 1, paragraph 4 Patent family members are listed in annex. Further documents are listed in the continuation of box C. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to "L" document which may throw doubts on priority claim(s) or involve an inventive step when the document is taken alone which is cited to establish the publication date of another document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the "O" document referring to an oral disclosure, use, exhibition or document is combined with one or more other such documents, such combination being obvious to a person skilled other means in the art. document published prior to the international filing date but "&" document member of the same patent family later than the priority date claimed Date of mailing of the international search report Date of the actual completion of the international search 27 -04- 1994 25 March 1994 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2

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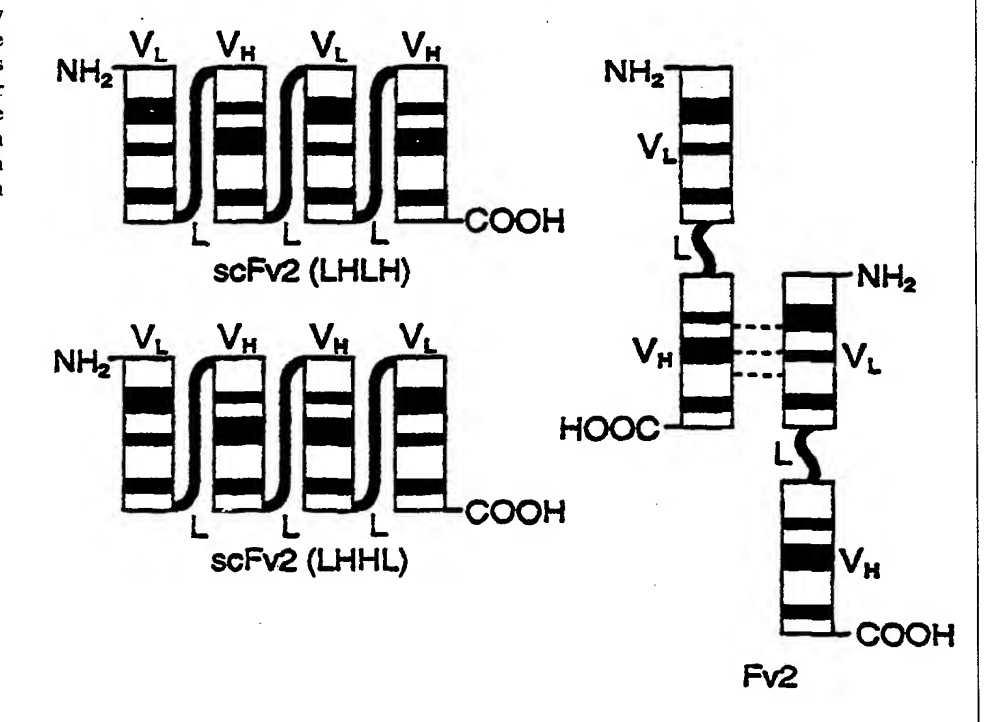
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### (54) Title: MULTIVALENT SINGLE CHAIN ANTIBODIES

### (57) Abstract

The present invention discloses multivalent single chain antibodies which have two or more biologically active antigen binding sites. multivalent single chain antibodies are formed by using a peptide linker to covalently link two or more single chain antibodies, each single chain antibody having a variable light domain linked to a variable heavy chain domain by a peptide linker.

### Schematic Representation Of Covalently & Non-Covalently Linked Single Chain Fv Multimers



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### **MULTIVALENT SINGLE CHAIN ANTIBODIES**

The present invention relates to single chain multivalent antibodies.

Antibodies are proteins belonging to a group of immunoglobulins elicited by the immune system in response to a specific antigen or substance which the body deems foreign. There are five classes of human antibodies, each class having the same basic structure. The basic structure of an antibody is a tetramer, or a multiple thereof, composed of two identical 'heterodimers each consisting of a light and a heavy chain. The light chain is composed of one variable (V) and one constant (C) domain, while a heavy chain is composed of one variable and three or more constant domains. The variable domains from both the light and heavy chain, designated V<sub>L</sub> and V<sub>H</sub> respectively, determine the specificity of an immunoglobulin, while the constant (C) domains carry out various effector functions.

Amino acid sequence data indicate that each variable domain comprises three complementarity determining regions (CDR) flanked by four relatively conserved framework regions (FR). The FR are thought to maintain the structural integrity of the variable region domain. The CDR have been assumed to be responsible for the binding specificity of individual antibodies and to account for the diversity of binding of antibodies.

As the basic structure of an antibody contains two heterodimers, antibodies are multivalent molecules. For example, the IgG classes have two identical antigen binding sites, while the pentameric IgM class has 10 identical binding sites.

Monoclonal antibodies having identical genetic parentage and binding specificity have been useful both as diagnostic and therapeutic agents. Monoclonal antibodies are routinely produced by hybridomas generated by fusion of mouse lymphoid cells with an appropriate mouse myeloma cell line according to established procedures. The administration of murine antibodies for in vivo therapy and diagnostics in humans is limited however, due to the human anti-mouse antibody response illicited by the human immune system.

Chimeric antibodies, in which the binding or variable regions of antibodies derived from one species are combined with the constant regions of antibodies derived from a different species, have been produced by recombinant DNA methodology. See, for example, Sahagen et al., *J. Immunol.*, 137:1066-1074 (1986); Sun et al., *Proc. Natl. Acad. Sci. USA*, 82:214-218 (1987); Nishimura et al., *Cancer Res.*, 47:999-1005 (1987); and Lie et al. *Proc Natl. Acad. Sci. USA*, 84:3439-3443 (1987) which disclose chimeric antibodies to tumor-associated antigens. Typically, the variable region of a murine antibody is joined with the constant region of a human antibody. It is expected that as such chimeric antibodies are largely human in composition, they will be substantially less immunogenic than murine antibodies.

Chimeric antibodies still carry the Fc regions which are not necessary for antigen binding, but constitute a major portion of the overall antibody structure which affects its pharmacokinetics. For the use of antibodies in immunotherapy or immunodiagnostics, is it

desirable to have antibody-like molecules which localize and bind to the target tissue rapidly and for the unbound material to quickly clear from the body. Generally, smaller antibody fragments have greater capillary permeability and are more rapidly cleared from the body than whole antibodies.

Since it is the variable regions of light and heavy chains that interact with an antigen, single chain antibody fragments (scFvs) have been created with one  $V_L$  and one  $V_H$ , containing all six CDR's, joined by a peptide linker (U.S. Patent 4,946,778) to create a  $V_L$ -L- $V_H$  polypeptide, wherein the L stands for the peptide linker. A scFv wherein the  $V_L$  and  $V_H$  domains are orientated  $V_H$ -L- $V_L$  is disclosed in U.S. Patent 5,132,405.

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As the scFvs have one binding site as compared to the minimum of two for complete antibodies, the scFvs have reduced avidity as compared to the antibody containing two or more binding sites.

than one binding site to enhance the avidity of the polypeptide, and retain or increase their antigen recognition properties. In addition, it would be beneficial to obtain multivalent scFvs which are bispecific to allow for recognition of different epitopes on the target tissue, to allow for antibody-based recruitment of other immune effector functions, or allow antibody capture of a therapeutic or diagnostic moiety.

It has been found that single chain antibody fragments, each having one  $V_H$  and one  $V_L$  domain covalently linked by a first peptide linker, can be covalently linked by a second peptide linker to form a multivalent single chain antibody which maintains the binding affinity of a whole antibody. In one embodiment, the present invention is a multivalent single chain antibody having affinity for an antigen wherein the multivalent single chain antibody comprises two or more light chain variable domains and two or more heavy chain variable domains; wherein, each variable domain is linked to at least one other variable domain.

In another embodiment, the present invention is a multivalent single chain antibody which comprises two or more single chain antibody fragments, each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:

- (a) a first polypeptide comprising a light chain variable domain;
- (b) a second polypeptide comprising a heavy chain variable domain; and
- (c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.

In another embodiment, the invention provides a DNA sequence which codes for a multivalent single chain antibody, the multivalent single chain antibody comprising two or more single chain antibody fragments, each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:

(a) a first polypeptide comprising a light chain variable domain;

(b) a second polypeptide comprising a heavy chain variable domain; and

(c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.

The multivalent single chain antibodies allow for the construction of an antibody fragment which has the specificity and avidity of a whole antibody but are smaller in size allowing for more rapid capillary permeability. Multivalent single chain antibodies also allow for the construction of a multivalent single chain antibody wherein the binding sites can be two different antigenic determinants.

### BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 illustrates covalently linked single chain antibodies having the configuration  $V_L-L-V_H-L-V_H$  (LHLH) and  $V_L-L-V_H-L-V_H-L-V_L$  (LHHL) and a noncovalently linked Fv single chain antibody (Fv2).

Figure 2 illustrates the nucleotide sequence of CC49  $V_{L}$ .

Figure 3 illustrates the amino acid sequence of CC49 V<sub>L</sub>.

Figure 4 illustrates the nucleotide sequence of CC49 V<sub>H</sub>.

Figure 5 illustrates the amino acid sequence of CC49 V<sub>H</sub>.

Figure 6 illustrates the nucleotide sequence and amino acid sequence of the CC49 single chain antibody LHLH in p49LHLH.

Figure 7 illustrates the nucleotide sequence and amino acid sequence of the CC49  $_{
m O}$  single antibody LHHL in p49LHHL.

Figure 8 illustrates construction of plasmids pSL301 T and pSL301 HT.

Figure 9 illustrates construction of plasmid p49LHHL.

Figure 10 illustrates construction of plasmid p49LHLH.

Figure 11 illustrates the results of a competition assay using CC49 IgG, CC49 scFv2, and CC49 scFv using biotinylated CC49 IgG as competitor.

The entire teaching of all references cited herein are hereby incorporated by reference.

Nucleic acids, amino acids, peptides, protective groups, active groups and such, when abbreviated, are abbreviated according to the IUPAC IUB (Commission on Biological Nomenclature) or the practice in the fields concerned.

The term "single chain antibody fragment" (scFv) or "antibody fragment" as used herein means a polypeptide containing a  $V_L$  domain linked to a  $V_H$  domain by a peptide linker (L), represented by  $V_L$ -L- $V_H$ . The order of the  $V_L$  and  $V_H$  domains can be reversed to obtain polypeptides represented as  $V_H$ -L- $V_L$ . "Domain" is a segment of protein that assumes a discrete function, such as antigen binding or antigen recognition.

A "multivalent single chain antibody" means two or more single chain antibody fragments covalently linked by a peptide linker. The antibody fragments can be joined to form bivalent single chain antibodies having the order of the  $V_L$  and  $V_H$  domains as follows:

 $V_L-L-V_H-L-V_L-L-V_H$ ;  $V_L-L-V_H-L-V_L$ ;  $V_H-L-V_L-L-V_H-L-V_L$ ; or  $V_H-L-V_L-L-V_L-L-V_L-L-V_L$ . Single chain multivalent antibodies which are trivalent and greater have one or more antibody fragments joined to a bivalent single chain antibody by an additional interpeptide linker. In a preferred embodiment, the number of  $V_L$  and  $V_H$  domains is equivalent.

The present invention also provides for multivalent single chain antibodies which can be designated  $V_H$ -L- $V_H$ -L- $V_L$ -L- $V_L$ -L- $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_H$ .

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Covalently linked single chain antibodies having the configuration  $V_L-L-V_H-L-V_L-L-V_H$  (LHLH) and  $V_L-L-V_H-L-V_H-L-V_L$  (LHHL) are illustrated in Figure 1. A noncovalently linked Fv single chain antibody (Fv2) is also illustrated in Figure 1.

The single chain antibody fragments for use in the present invention can be derived from the light and/or heavy chain variable domains of any antibody. Preferably, the light and heavy chain variable domains are specific for the same antigen. The individual antibody fragments which are joined to form a multivalent single chain antibody may be directed against the same antigen or can be directed against different antigens.

To prepare a vector containing the DNA sequence for a single chain multivalent antibody, a source of the genes encoding for these regions is required. The appropriate DNA sequence can be obtained from published sources or can be obtained by standard procedures known in the art. For example, Kabat et al., Sequences of Proteins of Immunological Interest 4th ed., (1991), published by The U.S. Department of Health and Human Services, discloses sequences of most of the antibody variable regions which have been described to date.

When the genetic sequence is unknown, it is generally possible to utilize cDNA sequences obtained from mRNA by reverse transcriptase mediated synthesis as a source of DNA to clone into a vector. For antibodies, the source of mRNA can be obtained from a wide range of hybridomas. See, for example, the catalogue ATCC Cell Lines and Hybridomas, American Type Culture Collection, 20309 Parklawn Drive, Rockville Md., USA (1990). Hybridomas secreting monoclonal antibodies reactive with a wide variety of antigens are listed therein, are available from the collection, and usable in the present invention. These cell lines and others of similar nature can be utilized as a source of mRNA coding for the variable domains or to obtain antibody protein to determine amino acid sequence of the monoclonal antibody itself.

Variable regions of antibodies can also be derived by immunizing an appropriate vertebrate, normally a domestic animal, and most conveniently a mouse. The immunogen will be the antigen of interest, or where a hapten, an antigenic conjugate of the hapten to an antigen such as keyhole limpet hemocyanin (KLH). The immunization may be carried out conventionally with one or more repeated injections of the immunogen into the host mammal, normally at two to three week intervals. Usually, three days after the last challenge, the spleen is removed and dissociated into single cells to be used for cell fusion to provide hybridomas from which mRNA can readily be obtained by standard procedures known in the art.

When an antibody of interest is obtained, and only its amino acid sequence is known, it is possible to reverse translate the sequence.

The  $V_L$  and  $V_H$  domains for use in the present invention are preferably obtained from one of a series of CC antibodies against tumor-associated glycoprotein 72 antigen (TAG-72) disclosed in published PCT Application WO 90/04410 on May 3, 1990, and published PCT Application WO 89/00692 on January 26, 1989. More preferred are the  $V_L$  and  $V_H$  domains from the monoclonal antibody designated CC49 in PCT Publications WO 90/04410 and WO 89/00692. The nucleotide sequence (SEQ ID NO: 1) which codes for the  $V_L$  of CC49 is substantially the same as that given in Figure 1. The amino acid sequence (SEQ ID NO: 2) of the  $V_L$  of CC49 is substantially the same as that given in Figure 2. The nucleotide sequence (SEQ ID NO: 3) which codes for the  $V_H$  of CC49 is substantially the same as that given in Figure 3. The amino acid sequence (SEQ ID NO: 4) for the  $V_H$  of CC49 is substantially the same as that given in Figure 4.

present invention, it is necessary to have a suitable peptide linker. Suitable linkers for joining the V<sub>H</sub> and V<sub>L</sub> domains are those which allow the V<sub>H</sub> and V<sub>L</sub> domains to fold into a single polypeptide chain which will have a three dimensional structure very similar to the original structure of a whole antibody and thus maintain the binding specificity of the whole antibody from which antibody fragment is derived. Suitable linkers for linking the scFvs are those which allow the linking of two or more scFvs such that the V<sub>H</sub> and V<sub>L</sub> domains of each immunoglobulin fragment have a three dimensional structure such that each fragment maintains the binding specificity of the whole antibody from which the immunoglobulin fragment is derived. Linkers having the desired properties can be obtained by the method disclosed in U.S. Patent 4,946,778, the disclosure of which is hereby incorporated by reference.

From the polypeptide sequences generated by the methods described in the 4,946,778, genetic sequences coding for the polypeptide can be obtained.

Preferably, the peptide linker joining the  $V_H$  and  $V_L$  domains to form a scFv and the peptide linker joining two or more scFvs to form a multivalent single chain antibody have substantially the same amino acid sequence.

It is also necessary that the linker peptides be attached to the antibody fragments such that the binding of the linker to the individual antibody fragments does not interfere with the binding capacity of the antigen recognition site.

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A preferred linker is based on the helical linker designated 205C as disclosed in Pantoliano et al. *Biochem.*, 30, 10117-10125 (1991) but with the first and last amino acids changed because of the codon dictated by the Xho I site at one end and the Hind III site at the other. The amino acid sequence (SEQ ID NO: 5) of the preferred linker is as follows:

The linker is generally 10 to 50 amino acid residues. Preferably, the linker is 10 to 30 amino acid residues. More preferably the linker is 12 to 30 amino acid residues. Most preferred is a linker of 15 to 25 amino acid residues.

Expression vehicles for production of the molecules of the invention include plasmids or other vectors. In general, such vectors contain replicon and control sequences which are derived from species compatible with a host cell. The vector ordinarily carries a replicon site, as well as specific genes which are capable of providing phenotypic selection in transformed cells. For example, *E. coli* is readily transformed using pBR322 [Bolivar et al., *Gene*, 2, 95- (1977), or Sambrook et al., *Molecular Cloning*, Cold Spring Harbor Press, New York, 2nd Ed. (1989)].

Plasmids suitable for eukaryotic cells may also be used. *S. cerevisiae*, or common baker's yeast, is the most commonly used among eukaryotic microorganisms, although a number of other strains, such as Pichia pastoris, are available. Cultures of cells derived from multicellular organisms such as SP2/0 or Chinese Hamster Ovary (CHO), which are available from the ATCC, may also be used as hosts. Typical of vector plasmids suitable for mammalian cells are pSV2neo and pSV2gpt (ATCC); pSVL and pKSV-10 (Pharmacia), pBPV-1/pML2d (International Biotechnology, Inc.).

The use of prokaryotic and eukaryotic viral expression vectors to express the genes for polypeptides of the present invention is also contemplated.

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It is preferred that the expression vectors and the inserts which code for the single chain multivalent antibodies have compatible restriction sites at the insertion junctions and that those restriction sites are unique to the areas of insertion. Both vector and insert are treated with restriction endonucleases and then ligated by any of a variety of methods such as those described in Sambrook et al., supra.

Preferred genetic constructions of vectors for production of single chain multivalent antibodies of the present invention are those which contain a constitutively active transcriptional promoter, a region encoding signal peptide which will direct synthesis/secretion of the nascent single chain polypeptide out of the cell. Preferably, the expression rate is commensurate with the transport, folding and assembly steps to avoid accumulation of the polypeptide as insoluble material. In addition to the replicon and control sequences, additional elements may also be needed for optimal synthesis of single chain polypeptide. These elements may include splice signals, as well as transcription promoter, enhancers, and termination signals. Furthermore, additional genes and their products may be required to facilitate assembly and folding (chaperones).

Vectors which are commercially available can easily be altered to meet the above criteria for a vector. Such alterations are easily performed by those of ordinary skill in the art in light of the available literature and the teachings herein.

Additionally, it is preferred that the cloning vector contain a selectable marker, such as a drug resistance marker or other marker which causes expression of a selectable trait by the host cell. "Host cell" refers to cells which can be recombinantly transformed with vectors constructed using recombinant DNA techniques. A drug resistance or other selectable marker is intended in part to facilitate in the selection of transformants. Additionally, the presence of a selectable marker, such as a drug resistance marker, may be of use in keeping contaminating microorganisms from multiplying in the culture medium. In this embodiment, such a pure culture of the transformed host cell would be obtained by culturing the cells under conditions which require the induced phenotype for survival.

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Recovery and purification of the present invention can be accomplished using standard techniques known in the art. For example, if they are secreted into the culture medium, the single chain multivalent antibodies can be concentrated by ultrafiltration. When the polypeptides are transported to the periplasmic space of a host cell, purification can be accomplished by osmotically shocking the cells, and proceeding with ultrafiltration, antigen affinity column chromatography or column chromatography using ion exchange chromatography and gel filtration. Polypeptides which are insoluble and present as refractile bodies, also called inclusion bodies, can be purified by lysis of the cells, repeated centrifugation and washing to isolate the inclusion bodies, solubilization, such as with guanidine-HCl, and refolding followed by purification of the biologically active molecules.

The activity of single chain multivalent antibodies can be measured by standard assays known in the art, for example competition assays, enzyme-linked immunosorbant assay (ELISA), and radioimmunoassay (RIA).

The multivalent single chain antibodies of the present invention provide unique benefits for use in diagnostics and therapeutics. The use of multivalent single chain antibodies afford a number of advantages over the use of larger fragments or entire antibody molecules. They reach their target tissue more rapidly, and are cleared more quickly from the body.

For diagnostic and/or therapeutic uses, the multivalent single chain antibodies can be constructed such that one or more antibody fragments are directed against a target tissue and one or more antibody fragments are directed against a diagnostic or therapeutic agent.

The invention also concerns pharmaceutical compositions which are particularly advantageous for use in the diagnosis and/or therapy of diseases, such as cancer, where target antigens are often expressed on the surface of cells. For diagnostic and/or therapeutic uses, the multivalent single chain antibodies can be conjugated with an appropriate imaging or therapeutic agent by methods known in the art. The pharmaceutical compositions of the invention are prepared by methods known in the art, e.g., by conventional mixing, dissolving or lyophilizing processes.

The invention will be further clarified by a consideration of the following examples, which are intended to be purely exemplary of the present invention.

### **ABBREVIATIONS**

	BCIP	5-bromo-4-chloro-3-indoyl phosphate
	bp	base pair
5	Bis-Tris propane	<pre>(1,3-bis[tris(hydroxymethyl)-methylamino]- propane)</pre>
	BSA	bovine serum albumin
	CDR	Complementarity determining region
	ELISA	enzyme linked immunosorbent assay
	Fv2	non-covalent single chain Fv dimer
10	IEF	isoelectric focusing
	Kbp	kilo base pair
	LB	Luria-Bertani medium
	Mab	monoclonal antibody
	MES	2-(N-Morpholino)ethane sulfonic acid
15	MW	molecular weight
	NBT	nitro blue tetrazolium chloride
	Oligo	Oligonucleotides
	PAG	polyacrylamide gel
	PAGE	polyacrylamide gel electrophoresis
20	PBS	phosphate buffered saline
·	PCR	polymerase chain reaction
	pSCFV	plasmid containing DNA sequence coding for SCFV
	RIGS	radioimmunoguided surgery
25	RIT	radioimmunotherapy
	scFv	single chain Fv immunoglobulin fragment monomer
	scFv2	single chain Fv immunoglobulin fragment dimer covalently linked
	SDS	sodium dodecyl sulfate
	TBS	Tris-buffered saline
30	Ťris	(Tris[hydroxymethyl]aminomethane)
	TTBS	Tween-20 wash solution
	VH	immunoglobulin heavy chain variable domain
	V <sub>L</sub>	immunoglobulin light chain variable domain

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### **Antibodies**

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CC49: A murine monoclonal antibody specific to the human tumor-associated glycoprotein 72 (TAG-72) deposited as ATCC No. HB9459.

CC49 FAB: An antigen binding portion of CC49 consisting of an intact light chain linked to the N-terminal portion of the heavy chain.

<u>CC49 scFv</u>: Single chain antibody fragment consisting of two variable domains of CC49 antibody joined by a peptide linker.

CC49 Fv2: Two CC49 scFv non-covalently linked to form a dimer. The number after Fv refers to the number of monomer subunits of a given molecule, e.g., CC49 Fv6 refers to the hexamer multimers.

CC49 scFv2: Covalently-linked single chain antibody fragment consisting of two CC49  $V_L$  domains and two  $V_H$  domains joined by three linkers. Six possible combinations for the order of linking the  $V_L(L)$  and the  $V_H(H)$  domains together are: LHLH, LHHL, LLHH, HLLH, and HHLL.

### 15 Plasmids

pSCFV UHM: Plasmid containing coding sequence for scFv consisting of a CC49 variable light chain and a CC49 variable heavy chain joined by a 25 amino acid linker.

p49LHLH or p49LHHL: Plasmids containing the coding sequence for producing CC49 scFv2 LHLH or LHHL products, respectively.

### 20 EXAMPLES

### **General Experimental**

Procedures for molecular cloning are as those described in Sambrook et al., Molecular Cloning, Cold Spring Harbor Press, New York, 2nd Ed. (1989) and Ausubel et al., Current Protocols in Molecular Biology, John Wiley and Sons, New York (1992), the disclosures of which are hereby incorporated by reference.

All water used throughout was deionized distilled water.

### Oligonucleotide Synthesis and Purification

All oligonuclotides (oligos) were synthesized on either a Model 380A or a Model 391 DNA Synthesizer from Applied Biosystems (Foster City, CA) using standard β-cyanoethyl phosphoramidites and synthesis columns. Protecting groups on the product were removed by heating in concentrated ammonium hydroxide at 55°C for 6 to 15 hours. The ammonium hydroxide was removed through evaporation and the crude mixtures were resuspended in 30 to 40 μL of sterile water. After electrophoresis on polyacrylamide-urea gels, the oligos were visualized using short wavelength ultraviolet (UV) light. DNA bands were excised from the gel and eluted into 1 mL of 100 mM Tris-HCl, pH 7.4, 500 mM NaCl, 5 mM EDTA over 2 hours at 65°C. Final purification was achieved by applying the DNA to Sep-Pac C-18 columns (Millipore, Bedford, MA) and eluting the bound oligos with 60 percent methanol. The

solution volume was reduced to approximately 50  $\mu$ L and the DNA concentration was determined by measuring the optical density at 260 nm (OD<sub>260</sub>).

### **Restriction Enzyme Digests**

All restriction enzyme digests were performed using Bethesda Research
Laboratories (Gaithersburg, MD), New England Biolabs, Inc. (Beverly, MA) or Boehringer
Mannheim (BM, Indianapolis, IN) enzymes and buffers following the manufacturer's
recommended procedures. Digested products were separated by polyacrylamide gel
electrophoresis (PAGE). The gels were stained with ethidium bromide, the DNA bands were
visualized using long wavelength UV light and the DNA bands were then excised. The gel slices
were placed In dialysis tubing (Union Carbide Corp., Chicago) containing 5 mM Tris, 2.5 mM
acetic acid, 1 mM EDTA, pH 8.0 and eluted using a Max Submarine electrophoresis apparatus
(Hoefer Scientific Instruments, CA). Sample volumes were reduced on a Speed Vac
Concentrator (Savant Instruments, Inc., NY). The DNA was ethanol precipitated and redissolved
in sterile water.

### 5 Enzyme Linked Immunosorbent Assay (ELISA)

TAG-72 antigen, prepared substantially as described by Johnson et al, Can. Res., 46, 850-857 (1986), was adsorbed onto the wells of a polyvinyl chloride 96 well microtiter plate (Dynatech Laboratories, Inc., Chantilly, VA) by drying overnight. The plate was blocked with 1 percent BSA in PBS for 1 hour at 31°C and then washed 3 times with 200 µL of PBS, 0.05 percent Tween-20. 25 µL of test antibodies and 25 µL of biotinylated CC49 (1/20,000 dilution of a 1 mg/mL solution) were added to the wells and the plate incubated for 30 minutes at 31°C. The relative amounts of TAG-72 bound to the plate, biotinylated CC49, streptavidinalkaline phosphatase, and color development times were determined empirically in order not to have excess of either antigen or biotinylated CC49, yet have enough signal to detect competition by scFv. Positive controls were CC49 at 5 µg/mL and CC49 Fab at 10 µg/mL. Negative controls were 1 percent BSA in PBS and/or concentrated LB. Unbound proteins were washed away. 50 µL of a 1:1000 dilution of streptavidin conjugated with alkaline phosphatase (Southern Biotechnology Associates, Inc., Birmingham, AL) were added and the plate was incubated for 30 minutes at 31°C. The plate was washed 3 more times. 50 µL of a para-nitrophenyl-phosphate solution (Kirkegaard & Perry Laboratories, Inc., Gaithersburg, MD) were added and the color reaction was allowed to develop for a minimum of 20 minutes. The relative amount of scFv2 binding was measured by optical density scanning at 404-450 nm using a microplate reader (Molecular Devices Corporation, Manlo Park, CA). Binding of the scFv2 species resulted in decreased binding of the biotinylated CC49 with a concomitant decrease in color development.

### **SDS-PAGE and Western Blotting**

Samples for SDS-PAGE analysis (20  $\mu$ L) were prepared by boiling in a non-reducing sample preparation buffer-Seprasol I (Integrated Separation Systems (ISS), Natick, MA) for

PCT/US93/12039 WO 94/13806

5 minutes and loaded on 10-20 percent gradient polyacrylamide Daiichi Minigels as per the manufacturer's directions (ISS).

Electrophoresis was conducted using a Mini 2-gel apparatus (ISS) at 55 mA per gel at constant current for approximately 75 minutes. Gels were stained in Coomassie Brilliant Blue R-250 (Bio-Rad, Richmond, CA) for at least 1 hour and destained. Molecular weight standards were prestained (Mid Range Kit, Diversified Biotech, Newton Center, MA) and included the following proteins: Phosphorylase b, glutamate dehydrogenase, ovalbumin, lactate dehydrogenase, carbonic amhydrase, B-lactoglobulin and cytochrome C. The corresponding MWs are: 95,500, 55,000, 43,000, 36,000, 29,000, 18,400, and 12,400, respectively.

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When Western analyses were conducted, a duplicate gel was also run. After electrophoresis, one of the gels was equilibrated for 15-20 minutes in anode buffer #1 (0.3 M Tris-HCl pH 10.4). An Immobilion-P PVDF (polyvinylidene dichlorine) membrane (Millipore, Bedford, MA) was treated with methanol for 2 seconds, and immersed in water for 2 minutes. The membrane was then equilibrated in anode buffer #1 for 3 minutes. A Milliblot-SDE apparatus (Millipore) was utilized to transfer proteins in the gel to the membrane. A drop of anode buffer #1 was placed in the middle of the anode electrode surface. A sheet of Whatman 3MM filter paper was soaked in anode buffer #1 and smoothly placed on the electrode surface. Another filter paper soaked in anode buffer #2 (25 mM tris pH 10.4) was placed on top of the first one. A sandwich was made by next adding the wetted PVDF membrane, placing the equilibrated gel on top of this and finally adding a sheet of filter paper soaked in cathode buffer (25mM Tris-HCl, pH 9.4 in 40 mM glycine). Transfer was accomplished in 30 minutes using 250 mA constant current (initial voltage ranged from 8-20 volts).

After blotting, the membrane was rinsed briefly in water and placed in a dish with 20 mL blocking solution (1 percent bovine serum albumin (BSA) (Sigma, St. Louis, MO) in Tris-buffered saline (TBS)). TBS was purchased from Pierce Chemical (Rockford, IL) as a preweighed powder such that when 500 mL water is added, the mixture gives a 25 mM Tris, 0.15 M sodium chloride solution at pH 7.6. The membranes were blocked for a minimum of 1 hour at ambient temperature and then washed 3 times for 5 minutes each using 20 mL 0.5 percent Tween-20 wash solution (TTBS). To prepare the TTBS, 0.5mL of Tween 20 (Sigma) was mixed per liter of TBS. The probe antibody used was 20 mL biotinylated FAID14 solution (10 µg per 20 mL antibody buffer). Antibody buffer was made by adding 1 g BSA per 100 mL of TTBS. After probing for 30-60 minutes at ambient temperature, the membrane was washed 3 times with TTBS, as above.

Next, the membrane was incubated for 30-60 minutes at ambient temperature with 20 mL of a 1:500 dilution in antibody buffer of streptavidin conjugated with alkaline phosphatase (Southern Biotechnology Associates, Birmingham, AL). The wash step was again repeated after this, as above. Prior to the color reaction, membranes were washed for 2 minutes in an alkaline carbonate buffer (20 mL). This buffer is 0.1 M sodium bicarbonate,

1 mM MgCl<sub>2</sub>·H<sub>2</sub>0, pH 9.8. To make up the substrate for alkaline phosphatase, nitroblue tetrazolium (NBT) chloride (50 mg, Sigma) was dissolved in 70 percent dimethylformamide. 5-Bromo-4-chloro-3-indoyl phosphate (BClP) (25 mg, Sigma) was separately dissolved in 100 percent dimethylformamide. 5-Bromo-4-chloro-3-indoyl phosphate (BClP) 25 mg, Sigma) was separately dissolved in 100 percent dimethylformamide. These solutions are also commercially available as a Western developing agent sold by Promega. For color development, 120 µL of each were added to the alkaline solution above and allowed to react for 15 minutes before they were washed from the developed membranes with water.

**Biotinylated FAID14** 

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FAID14 is a murine anti-idiotypic antibody (IgG2a, K isotype) deposited as ATCC No. CRL 10256 directed against CC49. FAID14 was purified using a Nygene Protein A affinity column (Yonkers, NY). The manufacturer's protocol was followed, except that 0.1 M sodium citrate, pH 3.0 was used as the elution buffer. Fractions were neutralized to pH  $\sim$ 7 using 1.0 M Tris-HCl pH 9.0. The biotinylation reaction was set up as follows. FAID14 (1 mg, 100  $\mu$ L in water) was mixed with 100  $\mu$ L of 0.1 M Na<sub>2</sub>CO<sub>3</sub> pH 9.6. Biotinyl- $\epsilon$ -amino-caproic acid N-hydroxy succinimide ester (Biotin-X-NHS) (Calbiochem, LaJolla, CA) (2.5 mg) was dissolved in 0.5 mL dimethylsulfoxide. Biotin-X-NHS solution (20  $\mu$ L) was added to the FAID14 solution and allowed to react at 22°C for 4 hours. Excess biotin and impurities were removed by gel filtration, using a Pharmacia Superose 12 HR10/30 column (Piscataway, NJ). At a flow rate of 0.8 mL/min, the biotinylated FAID14 emerged with a peak at 16.8 min. The fractions making up this peak were pooled and stored at 4°C and used to detect the CC49 idiotype as determined by the CC49 V<sub>L</sub> and V<sub>R</sub> CDRs.

### Isoelectric Focusing (IEF)

Isoelectric points (pl's) were predicted using a computer program called PROTEIN--TITRATE, available through DNASTAR (Madison, WI). Based on amino acid composition with an input sequence, a MW value is given, in addition to the pl. Since Cys residues contribute to the charge, the count was adjusted to 0 for Cys, since they are all involved in disulfide bonds.

Experimentally, pl's were determined using Isogel agarose IEF plates, pH range 3-10 (FMC Bioproducts, Rockland, ME). A Biorad Bio-phoresis horizontal electrophoresis cell was used to run the IEF, following the directions of both manufacturers. The electrophoresis conditions were: 500 volts (limiting), at 20 mA current and 10 W of constant power. Focusing was complete in 90 min. IEF standards were purchased from Biorad; the kit included phycocyanin, β-lactoglobulin B, bovine carbonic anhydrase, human carbonic anhydrase, equine myoglobin, human hemoglobins A and C, 3 lentil lectins and cytochrome C, with pl values of 4.65, 5.10, 6.00, 6.50, 7.00, 7.10 and 7.50, 7.80, 8.00, and 8.20 and 9.60, respectively. Gels were stained and destained according to the directions provided by FMC.

### Quantitation of CC49 Antibody Species

All purified CC49 antibodies including the IgG, scFv2 species and the monomeric scFv were quantitated by measuring the absorbence of protein dilutions at 280 mm using matching 1.0 cm pathlength quartz cuvettes (Hellma) and a Perkin-Elmer UV/VIS Spectrophotometer, Model 552A. Molar absorptivities (E<sub>m</sub>) were determined for each antibody by using the following formula:

 $E_{m} = (number Trp) X 5,500 + (number Tyr) X 1,340 + (number (Cys)2) X 150 + (number Phe) X 10$ 

The values are based on information given by D. B. Wetlaufer, Advances in Protein Chemistry, 17, 375-378).

### High Performance Liquid Chromatography

All high performance liquid chromatography (HPLC) was performed for CC49 scFv2 purification using an LKB HPLC system with titanium or teflon tubing throughout. The system consists of the Model 2150 HPLC pump, model 2152 controller, UV CORD SII model 2238 detection system set at an absorbence of 276 nm and the model 2211 SuperRac fraction collector.

### **PCR Generation of Subunits**

All polymerase chain reactions (PCR) were performed with a reaction mixture consisting of: 150 picograms (pg) plasmid target (pSCFVUHM); 100 pmoles primers; 1 µL Perkin-Elmer-Cetus (PEC, Norwalk, CT) Ampli-Taq polymerase; 16 µL of 10 mM dNTPs and 10 µL of 10X buffer both supplied in the PEC kit; and sufficient water to bring the volume to total volume to 100 µL. The PCR reactions were carried out essentially as described by the manufacturer. Reactions were done in a PEC 9600 thermocycler with 30 cycles of: denaturation of the DNA at 94°C for 20 to 45 sec, annealing from between 52 to 60°C for 0.5 to 1.5 min., and elongation at 72°C for 0.5 to 2.0 min. Oligonucleotide primers were synthesized on an Applied Biosystems (Foster City, CA) 380A or 391 DNA synthesizer and purified as above.

Ligation reactions using 100 ng of vector DNA and a corresponding 1:1 stoichiometric equivalent of insert DNA were performed using a Stratagene (La Jolla, CA) T4 DNA ligase kit following the manufacturer's directions. Ligation reactions (20 µL total volume) were initially incubated at 18°C and allowed to cool gradually overnight to 4°C.

Transformations

Transformations were performed utilizing 100 µL of Stratagene E. coli AG1 competent cells (Stratagene, La Jolla, CA) according to the directions provided by the manufacturer. DNA from the ligation reactions (1-5 µL) were used. After the transformation step, cells were allowed to recover for 1 hr in Luria broth (LB) at 37°C with continuous mixing and subsequently plated onto either 20 µg/mL chloramphenicol containing (CAM 20) Luria agar for pSCFVUHM, p49LHLH or p49LHHL or 100 µg/mL ampicillin (AMP 100) Luria agar plates

(LB-AMP 100) for clones containing the plasmid pSL301 or subsequent constructions derived from pSL301.

### Screening of E. coli Clones

Bacterial plasmids were isolated from LB broth culture containing the appropriate drug to maintain selection pressure using Promega (Madison, WI) Magic mini-prep plasmid preparation kits. The kit was used per the manufacturer's specifications.

### **Plasmid Constructions**

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Two plasmids, designated p49LHLH and p49LHHL, were constructed to produce multivalent single chain antibodies. The host cell containing p49LHLH produced a polypeptide which can be designated by  $V_L-L-V_H-L-V_L-L-V_H$  where  $V_L$  and  $V_H$  are the light and heavy cahin variable regions of CC49 antibody and linker (L) is a 25 amino acid linker having the sequence (SEQ ID NO: 5).

The host cell containing p49LHHL produced a polypeptide which can be designated by  $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_L$  where  $V_L$  and  $V_H$  are the light and heavy chain variable domains of the CC49 antibody and L is a peptide linker having the amino acid sequence indicated above.

The nucleotide sequence (SEQ ID NO: 6) and amino acid sequence (SEQ ID NO: 7) of the CC49  $V_L$ -L- $V_H$ -L- $V_L$ -L- $V_H$  (p49LHLH) are given in Figure 6. The nucleotide sequence (SEQ ID NO: 8) and amino acid sequence (SEQ ID NO: 9) of the CC49  $V_L$ -L- $V_H$ -L- $V_H$ -L- $V_L$  (p49LHHL) are given in Figure 7.

### Construction of pSL301 HT

penicillinase P (penP) terminator sequence was removed from the plasmid designated pSCFV UHM by a 45 minute digest with Nhe I and BamH I, excised from a 4.5 percent polyacrylamide gel after electrophoresis, electroeluted, ethanol precipitated and ligated into the same sites in the similarly prepared vector: pSL301 (Invitrogen, San Diego, CA). A procedure for preparing pSCFV UHM is given is U.S. patent application Ser. No. 07/935,695 filed August 21, 1992, the disclosure of which is hereby incorporated by reference. In general, pSCFV UHM contains a nucleotide sequence for a penP promoter; a unique Nco I restriction site; CC49 V<sub>L</sub> region; Hind III restriction site; a 25 amino acid linker; a unique a Xho I restriction site; CC49 V<sub>H</sub> region; Nhe I restriction site; penP terminator; and BamH I restriction site (see, Figure 8). The penP promoter and terminator are described in Mezes, et al. (1983), *J. Biol.* 

An aliquot of the ligation reaction (3  $\mu$ L) was used to transform competent *E. coli* AG1 cells which were plated on LB-AMP100 agar plates and grown overnight. Potential clones containing the penP terminator insert were screened using a Pharmacia (Gaithersburg, MD) T7

Quickprime <sup>32</sup>P DNA labeling kit in conjunction with the microwave colony lysis procedure outlined in Buluwela et al., *Nucleic Acid Research*, <u>17</u>, 452 (1989). The probe, which was the penP-Nhe I-BamH I terminator fragment itself was prepared and used according to the directions supplied with the Quickprime kit. A clone which was probe positive and which contained the 207 base pair inserts from a BamH I and Nhe I digest (base pairs (bp) 1958 to 2165, Figure 6) was designated pSL301 T and chosen to construct pSL301 HT which would contain the nucleotide sequence for CC49 V<sub>H</sub>. The reason the Nhe I-BamH I penP terminator was placed into pSL301 was to eliminate the Eco47 III restriction endonuclease site present in the polylinker region between its Nhe I and BamH I sites. This was designed to accommodate the subsequent build-up of the V<sub>L</sub> and V<sub>H</sub> domains where the Eco47 III site needed to be unique for the placement of each successive V domain into the construction. As each V domain was added at the Eco47 III-Nhe I sites, the Eco47 III was destroyed in each case to make the next Eco47 III site coming in on the unique insert.

The V<sub>H</sub> sequence was made by PCR with oligos 5' SCP1 and 3'oligo SCP5 using pSCFV UHM as the target for PCR amplification. The DNA sequence for SCP1 (SEQ ID NO: 10) and SCP5 (SEQ ID NO: 11) are as follows:

SCP1: 5'-TAAA CTC GAG GTT CAG TTG CAG CAG -3'

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SCP5: 5'-TAAA GCT AGC ACCA AGC GCT TAG TGA GGA GAC GGT GAC TGA GGT-3'
The underlined portion indicates the endonuclease restriction sites.

The amplified  $V_H$  DNA was purified from a 4 percent PAG, electroeluted ethanol precipitated and dissolved in 20  $\mu$ L water. The  $V_H$  sequence was digested with Xho I and Nhe I restriction enzymes and used as the insert with the pSL301 T vector which had been digested with the same restriction enzymes and subsequently purified. A standard ligation reaction was done and an aliquot (4  $\mu$ L) used to transform competent E. coli AG1 cells. The transformed cells were plated onto LB AMP100 agar plates. Candidate clones were picked from a Nhe I and Xho I digest screen that revealed that the CC49V<sub>H</sub> insert had been obtained.

DNA sequencing was performed to verify the sequence of the CC49V<sub>H</sub> with United States Biochemical (USB) (Cleveland, Ohio) Sequence kit and sequencing primers pSL301SEQB (a 21 bp sequencing primer which annealed in the pSL301 vector 57 bp upstream from the Xho I site) and CC49VHP, revealed clones with the correct CC49V<sub>H</sub> sequence in pSL301HT. This plasmid was used as the starting point in the construction of both pSL301-HLHT and pSL301-HLHT. The sequencing oligos used are shown here.

The nucleotide sequence of pSL301SEQ B (SEQ ID NO: 12) and CC49V<sub>H</sub> (SEQ ID No:  $\frac{1}{13}$ ) are as follows:

pSL301SEQB: 5'-TCG TCC GAT TAG GCA AGC TTA-3'
CC49VHP: 5'-GAT GAT TTT AAA TAC AAT GAG-3'

PCT/US93/12039 WO 94/13806

### Example 1 p49LHHL Construction

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Using pSL301 HT (5 µg) as the starting material, it was digested with Eco47 III and Nhe I and the larger vector fragment was purified. A CC49V<sub>H</sub> insert fragment was generated by PCR using SCP6C as the 5' oligo and SCP5 as the 3' oligo. The nucleotide sequence (SEQ ID NO: 14) of SCP6B is as follows:

5'- TAAA TGC GCA GAT GAC GCA AAG AAA GAC GCA GCT AAA AAA GAC GAT SCP6B: GCC AAA AAG GAT GAC GCC AAG AAA GAT CTT GAG GTT CAG TTG CAG CAG TCT-G'

The oligo SCP6B also contains part of the coding region for the linker (bp 8-76 of SEQ ID NO: 14). The portion of the oligo designed to anneal with the CC49VH target in pSCFV UHM is from bp77-90 in SEQ ID NO: 14.

The underlined sequence corresponds to the Fsp I site. The resulting PCR insert was purified, digested with Fsp I and Nhe I and used in a ligation reaction with the pSL301 HT Eco47 III-Nhe I vector (Figure 7). Competent E. coli AG1 cells were used for the transformation of this ligation reaction (3 µL) and were plated on LB-AMP100 agar plates. Two clones having the correct size Xho I-Nhe I insert representative of the pSL301 HHT product were sequenced with the oligo SQP1 and a single clone with the correct sequence (nucleotides 1124-1543 of Figure 7) was chosen for further construction. The nucleotide sequence of SQP1 (SEQ ID NO: 16) is as follows:

SQP1: 5'-TG ACT TTA TGT AAG ATG ATG T-3'

The final linker-V<sub>L</sub> subunit (bp 1544-1963, Figure 7) was generated using the 5'oligo, SCP7b and the 3' oligo, SCP8a, using pSCFV UHM as the target for the PCR. The nucleotide sequence of SCP7b (SEQ ID NO: 17 is as follows:

SCP7b: 5'-TAAA TGC GCA GAT GAC GCA AAG AAA GAC GCA GCT AAA AAA GAC GAT GCC AAA AAG GAT GAC GCC AAG AAA GAT CTT GAC ATT GTG ATG TCA CAG TCT 25 CC

The underlined nucleotides correspond to an Fsp I site. The nucleotide sequence of SCP8a (SEQ ID NO: 18) is as follows:

> 5'-TAAA GCT AGC TTT TTA CTT AAG CAC CAG CTT GGT CCC-3' SCP8a:

The first set of underlined nucleotides correspond to an Nhe I site, while the other corresponds to an Afl II site. Nucleotides 8-76 of SCP70 code for the linker (nucleotides 1544-1612 of Figure 7) while nucleotides 77-99 which anneal to the  $V_L$  correspond to 1613-1635 of Figure 7. The primer SCP8a has a short tail at its 5' end, a Nhe I restriction site, a stop codon, an Afl II restriction site and the last 21 bases of the  $V_L$ . After Fsp I and Nhe I digestion, this resulting 420 bp insert was purified and ligated into the Nhe I and Eco47 III sites of the purified pSL301HHT vector, candidate clones were screened with Nhe I and Xho I, the correct size insert verified and sequenced with 49LFR2(-) and SQP1 to confirm the newly inserted sequence in pSL301HHLT. The nucleotide sequence (SEQ ID NO: 19) is as follows:

49LFR2(-): 5'-CTG CTG GTA CCA GGC CAA G-3'

The plasmid pSL301HHLT was digested with Xho I and Nhe I, purified, and the resulting 1179 bp V<sub>H</sub>-linker-V<sub>H</sub>-linker-V<sub>L</sub> segment ligated into pSCFV UHM, which had been cut with the same restriction enzymes and the larger vector fragment purified, to form p49LHHL. The ligation reaction (4 µL aliquot) was used to transform competent E. coli AG1 cells (Stratagene) and plated onto LBCAM20 agar plates. A single clone which had a plasmid with the correct restriction enzyme map was selected to contain p49LHHL. The p49LHHL contains a penP promoter and a nucleotide sequence for the CC49 multivalent single chain antibody scFv2:

 $V_L-L-V_H-L-V_H-L-V_L$  or CC49 scFv2 (LHHL).

Example 2: p49LHLH Construction

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The construction of p49LHLH is schematically represented in Figure 11. A linker- $V_L$  subunit was generated with the 5' oligo SCP7b and the 3'oligo SCP9.

SCP9: 5'-TAA AGC TAG CAC CAA GCG CTT AGT TTC AGC ACC AGC TTG GTC CCA G-3'

The SCP7b oligo (nucleotides 8-76) codes for the linker in Figure 6 (corresponding to nucleotides 1124-1192) and annealed to the pSCFV UHM target for the PCR (nucleotides 77-99) corresponding to nucleotides 1193-1215 of the V<sub>L</sub> in Figure 6.

SCP9 has a Nhe I site (first underlined nucleotides) and an Eco47 III site (second underlined nucleotides) which are restriction sites needed for making the pSL301HLT ready to accept the next V domain. Nucleotides 18-23 of SCP9 correspond to nucleotides 1532-1537 of Figure 6 (coding for the first 2 amino acids of the linker), while nucleotides 24-46 correspond to nucleotides 1508-1531 of Figure 6 which was also the annealing region for SCP9 in the PCR. The plasmid pSL301 HT was digested with Eco47 III and Nhe I and the larger vector fragment was purified for ligation with the linker-CC49V<sub>L</sub> DNA insert fragment from the PCR which had been treated with Fsp I and Nhe I and purified. The ligation mixture (3 µL) was used to transform *E. coli* AG1 competent cells and one colony having the correct Xho I-Nhe I size fragment was sequenced using the oligo PENPTSEQ2. The nucleotide sequence (SEQ. ID NO. 21) is as follows:

5'-TTG ATC ACC AAG TGA CTT TAT G-3'

The sequencing results indicated that there had been a PCR error and deletion in the resulting pSL301HT clone. A five base deletion, corresponding to nucleotides 1533-1537 as seen in Figure 6 had been obtained and nucleotide 1531 which should have been a T was actually a G, as determined from the DNA sequence data. The resulting sequence was

5'...G AAGC GCT T...etc.

where the underlined sequence fortuitously formed an Eco47 III site. The AGCGCT sequence in Figure 6, would correspond to nucleotides 1530, 1531, 1532, 1538, 1539 and 1540. This error was corrected in the next step, generating pSL301 HLHT, by incorporating the 5 base deletion at the end of oligo SCP6C.

### SCP6C: 5'-TAAGCGCTGATGATGCTAAGAAGGACGCCGCAAAAAA GGACGACGCAAAAAAAGATGATGCAAAAAAAGGATCTGG AGGTTCAGTTGCAGCAGTCTGAC-3'

The underlined sequence in SCP6c corresponds to an Eco47 III site. SCP6C was used as the 5' oligo, with SCP10 as the 3' oligo in a PCR to generate a linker CC49  $V_L$  segment. The nucleotide sequence (SEQ ID NO: 23) is as follows:

SCP10: 5'TTG TGC TAG CTT TTT ATG AGG AGA CGG TGA CTG AGG TT-3'

The underlined sequence in SCP10 corresponds to the Nhe I site found at nucleotides 1958-1963 in Figure 6. The PCR insert was digested this time only with Nhe I and purified. The vector (pSL301 HLT) was digested at the Eco47 III site (that had been formed) and Nhe I and purified. The insert and vector were ligated and an aliquot (3 µL) used to transform competent E. coli AG1 cells. This was plated on LB-AMP100 plates and candidate clones screened with Xho I and Nhe I. Three clones having the correct size DNA were obtained. Two of these clones were sequenced using the oligo 49VLCDR3(+) and SQP1. The nucleotide sequence (DWQ ID NO: 24 of 49VLCDR3(+) is as follows:

49VLCDR3(+):

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5'-CAG CAG TAT TAT AGC TAT-3'

One clone, with the correct sequence was obtained and the sequence from nucleotides 1533 to 1963 in Figure 6 were verified, giving a correct pSL301 HLHL clone.

To generate the final plasmid, p49LHLH for expression in E. coli, pSL301 HLHT (5 μg) was digested with Nhe I and Xho I, and the smaller insert fragment containing the V<sub>H</sub>-L-V<sub>L</sub>-L-V<sub>H</sub> sequence purified. It was ligated with the larger purified vector fragment from a digest of pSCFV UHM (5 μg) with Xho I and Nhe I. An aliquot of the ligation mix (4 μL) was used to transform competent E. coli AG1 cells. The transformation mix was plated on LB-CAM20 plates, and a representative clone for p49 LHLH was selected on the basis of a correct restriction enzyme map (see Figure 10) and biological activity toward TAG-72.

Example 3: Purification of CC49 scFv2 LHLH and LHHL Covalently Linked Dimers

For the purification of the CC49 covalently linked single chain dimers, (scFv2),

E. coli periplasmic fractions were prepared from 1.0 L overnight cultures of both p49LHLH and
p49LHHL. Briefly, the culture was divided into 4 X 250 mL portions and centrifuged at
5,000 rpm for 10 minutes in a Sorvall GS-3 rotor. The pelleted cells were washed and
resuspended in 100 mL each of 10 mM Tris-HCl pH 7.3 containing 30 mM NaCl. The cells were
again pelleted and washed with a total of 100 mL 30 mM Tris-HCl pH 7.3 and pooled into one
tube. To this, 100 mL of 30 mM Tris-HCl pH 7.3 containing 40 percent w/v sucrose and 2.0 mL of
10 mM EDTA pH 7.5 was added. The mixture was kept at room temperature, with occasional
shaking, for 10 minutes. The hypertonic cells were then pelleted as before. In the next step, the
shock, the pellet was quickly suspended in 20 mL ice cold 0.5 mM MgCl<sub>2</sub> and kept on ice for 10
minutes, with occasional shaking. The cells were pelleted as before and the supernatant

containing the *E. coli* periplasmic fraction was clarified further by filtration through a 0.2 µm Nalge (Rochester, NY) filter apparatus and concentrated in Amicon (Danvers, MA) Centriprep 30 and Centricon 30 devices to a volume of less than 1.0 mL.

The concentrated periplasmic shockates from either the p49LHLH or p49LHHL clones were injected onto a Pharmacia (Piscataway, NJ) Superdex 75 HR 10/30 HPLC column that had been equilibrated with PBS. At a flow rate of 0.5 mL/minute, the product of interest, as determined by competition ELISA, had emerged between 21 through 24 minutes. The active 'fractions were pooled, concentrated as before and dialyzed overnight using a system 500 Microdialyzer Unit (Pierce Chemical) against 20 mM Tris-HCl pH 7.6 with 3-4 changes of buffer and using an 8,000 MW cut-off membrane. The sample was injected on a Pharmacia Mono Q HR 5/5 anion exchange HPLC column. A gradient program using 20 mM Tris-HCl pH 7.6 as buffer A and the same solution plus 0.5 M NaCl as buffer B was employed at a flow rate of 1.5 mL/min. The products of interest in each case, as determined by competition ELISA, emerged from the column between 3 and 4 minutes. Analysis of the fractions at this point on duplicate SDS-PAGE gels, one stained with Coomassie Brilliant Blue R-250 and the other transferred for Western analysis (using biotinylated FAID 14 as the probe antibody) revealed a single band at the calculated molecular weight for the scFv2 (LHLH or LHHL) species at 58,239 daltons. The active fractions were in each case concentrated, dialysed against 50 mM MES pH 5.8 overnight and injected on a Pharmacia Mono S HR 5/5 cation exchange column. The two fractions of interest from this purification step, as determined by SDS-PAGE and ELISA, fractions 5 and 6, eluted just before the start of the gradient, so they had not actually bound to the column. Fractions 5 and 6 were consequently pooled for future purification.

A Mono Q column was again run on the active Mono S fractions but the buffer used was 20 mM Tris-HCl, pH 8.0 and the flow rate was decreased to 0.8 mL/minute. The products emerged without binding, but the impurity left over from the Mono S was slightly more held up, so that separation did occur between 5 and 6 minutes. After this run, the products were homogeneous and were saved for further characterization.

### **Isoelectric Focusing**

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The isoelectric points (pl) of the constructs was predicted using the DNASTAR (Madison, WI) computer program Protein-titrate. Based on amino acid composition, a MW and pl value was calculated.

Experimentally, pls were determined using FMC Bioproducts (Rockland, ME) lsogel IEF plates, pH range 3-10. A Biorad (Richmond, CA) electrophoresis unit was used to run the IEF, following the directions of both manufacturers. The electrophoresis conditions were as follows: 500 V (limiting) at 20 mA and at 10 W of constant power. Focusing was complete in 90 minutes. Biorad IEF standards included phycocyanin, beta lactoglobulin B, bovine carbonic anhydrase, human carbonic anhydrase, equine myoglobulin, human hemoglobins A and C, 3 lentil lectin, and cytochrome C with pI value of 4.65, 5.10, 6.00, 6,50, 7.00, 7.50, 7.8, 8.00, 8.20

WO 94/13806 PCT/US93/12039

and 9.6, respectively. Gels were stained and destained according to directions provided by FMC. The DNASTAR program predicted values of 8.1 for the pl for both scFv2 species. A single, homogeneous band for the pure products was observed on the gel at pl values for both at 6.9.

Purified CC49 antibodies such as the IgG, scFv2 (LHLH and LHHL) were quantitated by measuring the absorbence spectrophotometrically at 280 nm. Molar absorbtivity values,  $\epsilon_{\rm M}$ , were determined for each using the formula cited above by Wetlaufer.

Based on the amino acid composition, the E<sup>0.1%</sup> (280 nanometers) values for CC49 IgG, CC49 scFv2 LHLH, CC49 scFv2 LHHL and CC49 scFv were 1.49, 1.65, 1.65 and 1.71, respectively.

#### 10 Example 4

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Relative activities of the CC49 scFv2 species LHLH and LHHL, were compared with the IgG and a monomer scFv form with a FLAG peptide at the COOH terminus.

Percent competition was determined from the ELISA data by the following equation:

Zero competition - sample reading (OD405-450 nm)  $_{\times 100}$  zero competition - 100 percent competition

The "zero competition" value was determined by mixing (1:1) one percent BSA with the biotinylated CC49 (3 X 10-14 moles) while the 100 percent competition value was based on a 5 µg/mL sample of CC49 lgG mixed with the biotinylated CC49 lgG. The data are presented in Figure 11. Absorbence values for the samples were measured at 405 nm - 450 nm.

The average of triplicate readings was used. Initially samples (25 µL) were applied to the TAG-72 coated microliter plates at 1.0 X 10-10 moles of binding sites/mL. Biotinylated CC49 (4 µg/µL diluted 1:20,000 - used 25 µL) diluted the samples by a factor of 2. Serial dilutions (1:2) were performed. Both forms of the scFv2 are approximately equivalent to the lgG (see Figure 11). In a separate experiment, a CC49 scFv monomer was compared to a Fab fragment,

both of which are monovalent and these were also shown to be equivalent in their binding affinity for TAG-72. These results indicate that both forms of the covalently linked dimers have 2 fully functional antigen binding sites. This is the same increase in avidity as observed with the whole IgG, relative to a monomeric species.

These data also indicate that the scFv2 molecules, like their CC49 IgG parent are candidates for immunotherapeutic applications, but with the benefit of increased capillary permeability and more rapid biodistribution pharmacokinetics. The advantage should allow multiple injections of compounds of the present invention and give higher tumor: tissue ratios in immunotherapeutic treatment regimens for cancer treatment, relative to the existing IgG molecules.

Other embodiments of the invention will be apparent to those skilled in the art from a consideration of this specification or practice of the invention disclosed herein. It is

WO 94/13806 PCT/US93/12039

intended that the specification and examples be considered as exemplary only, with the true scope and spirit of the invention being indicated by the following claims.

WO 94/13806 PCT/US93/12039

1. A mutivalent single chain antibody which comprises two or more single chain antibody fragments each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:

- (a) a first polypeptide comprising a light chain variable domain;
- (b) a second polypeptide comprising a heavy chain variable domain; and
- (c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.
- 2. The multivalent single chain antibody of Claim 1 wherein the first peptide linker has the amino and sequence

Leu Ser Ala Asp Asp Ala Lys Lys Asp Ala Lys Lys Asp Asp Ala Lys Lys Asp Ala Lys Lys Asp Leu.

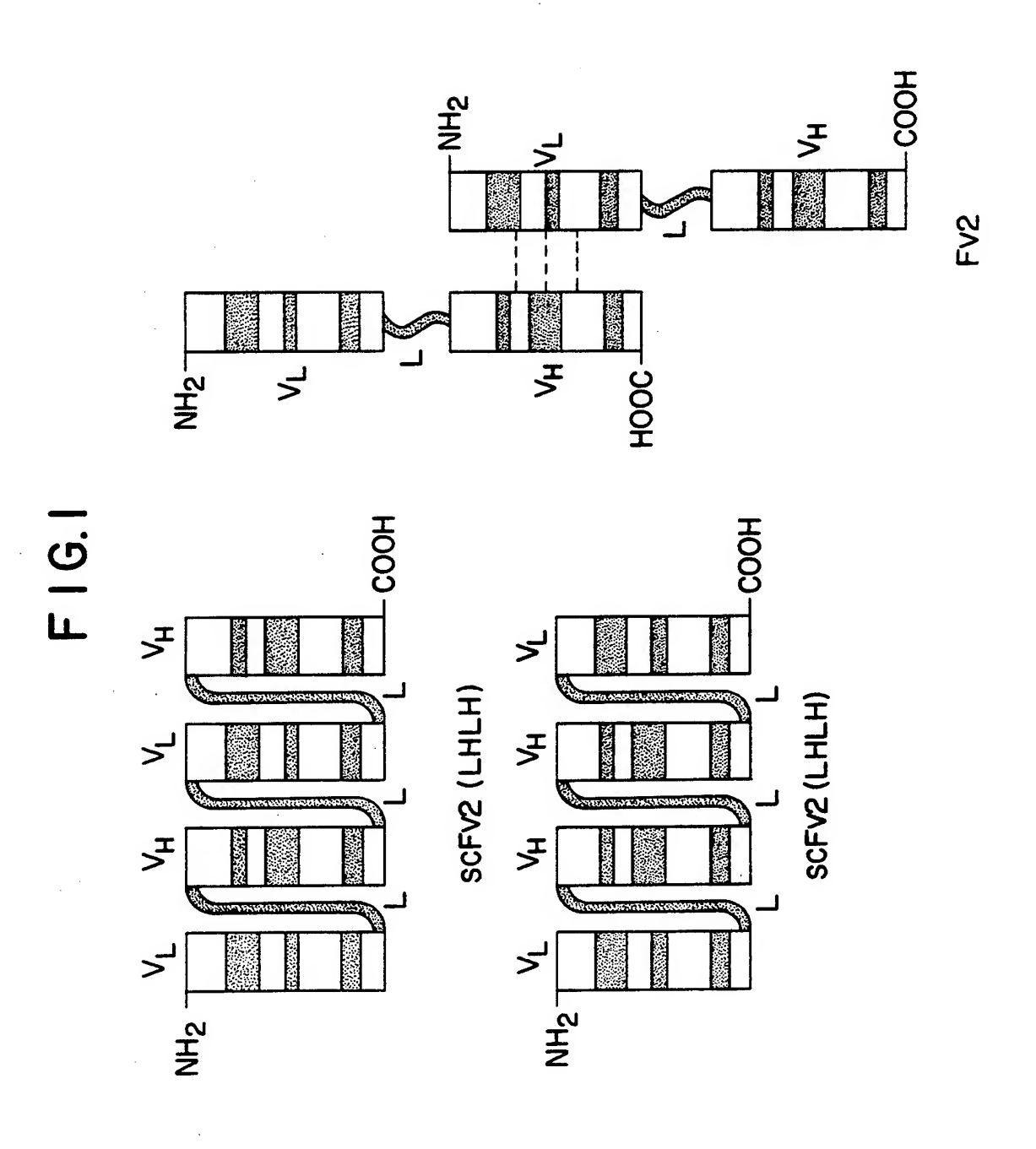
- 3. The multivalent single chain antibody of Claim 1 wherein the light chain variable region has an amino acid sequence substantially the same as that of Figure 3 and the heavy chain variable region has an amino acid sequence substantially the same as that of Figure 5.
- 4. The multivalent single chain antibody of Claim 1 wherein the first and second peptide linkers have an amino acid sequence which is substantially the same.
- 5. A DNA sequence which codes for a mutivalent single chain antibody, the multivalent single antibody comprising two or more single chain antibody fragments, each fragment having affinity for an antigen wherein the fragments are covalently linked by a first peptide linker and each fragment comprising:
  - (a) a first polypeptide comprising a light chain variable domain;
  - (b) a second polypeptide comprising a heavy chain variable domain; and
  - (c) a second peptide linker linking the first and second polypeptides into a functional binding moiety.
- 6. The DNA sequence of Claim 5 wherein the sequence coding for the first polypeptide is substantially the same as that of Figure 2 and the sequence coding for the second polypeptide is substantially the same as that of Figure 3.

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### FIG. 2

GAC ATT GTG ATG TCA CAG TCT CCA TCC TCC CTA CCT GTG TCA GTT GGC GAG AAG GTT ACT TTG AGC TGC AAG TCC AGT CAG AGC CTT TTA TAT AGT GGT AAT CAA AAG AAC TAC TTG GCC TGG TAC CAG CAG AAA CCA GGG CAG TCT CCT AAA CTG CTG ATT TAC TGG GCA TCC GCT AGG GAA TCT GGG GTC CCT GAT CGC TTC ACA GGC AGT GGA ACT GGA ACA GAT TTC ACT CTC TCC ATC AGC AGT GTG AAG ACT GAA GAC CTG GCA GTT TAT TAC TGT CAG CAG TAT TAT AGC TAT CCC CTC ACG TTC ACG CTG AAG

### FIG. 3

Asp Ile Val Met Ser Gln Ser Pro Ser Ser Leu Pro Val Ser Val Gly Glu Lys Val Thr Leu Ser Cys Lys Ser Ser Gln Ser Leu Leu Tyr Ser Gly Asn Gln Lys Asn Tyr Leu Ala Trp Tyr Gln Gln Lys Pro Gly Gln Ser Pro Lys Leu Leu Ile Tyr Trp Ala Ser Ala Arg Glu Ser Gly Val Pro Asp Arg Phe Thr Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Ser Ile Ser Ser Val Lys Thr Glu Asp Leu Ala Val Tyr Tyr Cys Gln Gln Tyr Tyr Ser Tyr Pro Leu Thr Phe Gly Ala Gly Thr Lys Leu Val Leu Lys

### FIG. 4

GAG GTT CAG TTG CAG CAG TCT GAC GCT GAG TTG GTG AAA CCT
GGG GCT TCA GTG AAG ATT TCC TGC AAG GCT TCT GGC TAC ACC
TTC ACT GAC CAT GCA ATT CAC TGG GTG AAA CAG AAC CCT GAA
CAG GGC CTG GAA TGG ATT GGA TAT TTT TCT CCC GGA AAT GAT
GAT TTT AAA TAC AAT GAG AGG TTC AAG GGC AAG GCC ACA CTG
ACT GCA GAC AAA TCC TCC AGC ACT GCC TAC GTG CAG CTC AAC
AGC CTG ACA TCT GAG GAT TCT GCA GTG TAT TTC TGT ACA AGA
TCC CTG AAT ATG GCC TAC TGG GGT CAA GGA ACC TCA GTC ACC
GTC TCC TCA

### FIG. 5

Glu Val Gln Leu Gln Gln Ser Asp Ala Glu Leu Val Lys Pro Gly Ala Ser Val Lys Ile Ser Cys Lys Ala Ser Gly Tyr Thr Phe Thr Asp His Ala Ile His Trp Val Lys Gln Asn Pro Glu Gln Gly Leu Glu Trp Ile Gly Tyr Phe Ser Pro Gly Asn Asp Asp Phe Lys Tyr Asn Glu Arg Phe Lys Gly Lys Ala Thr Leu Thr Ala Asp Lys Ser Ser Ser Thr Ala Tyr Val Gln Leu Asn Ser Leu Thr Ser Glu Asp Ser Ala Val Tyr Phe Cys Thr Arg Ser Leu Asn Met Ala Tyr Trp Gly Gln Gly Thr Ser Val Thr Val Ser Ser

382

Pro

430

Cys

Ser

Leu

Val

Lys

Glu GAG

Gly

Val

Ser

Val

Pro

Leu

Ser

Ser TCC

Thr ACT 478

A18 GCC

> Leu 116

Tyr

Asn

Lys

Glu

Asn

Gly

Ser

Tyr

Leac

Ser

Glu

Ser

Ser

30 Leu TTA

FIG. 61

သသ GAA CAA GAA TAC Leu TTA Ser CTT TCC GAC GTG TAC AGG Leu TTG Glu ACG ATA TCA 000 000 Gly Ser 17-T-NH-T-NH CCT AAA CCA TIC 999 Met ATG TII AAA TTT Ala GCT CGT R I ATT GTA TCA Ala GCC TTC Val GTG ACT GAT ECOR TGA A TCA AAA ATATATAT Ala GCA AAT ACT I1e ATT AGG CTG CGA AAC CAA R2-Thr TAT VL Asp GAC **6122** CAT AAT (PENPE PENPR1-AGC CAT ACG SSS ATA Pro Ala GCC QF OF GAA TAC Met ATG AGT Leu TAT ATC H SEQUENCE Nco Ala P GCC / GTG CCT TCT TIG Leu TCA ACG ACA ຽວວ AAA TGI Tyr Pro TCT GTT ACID TTG TCC CTG Gln CAA TGA TIT GAT Lys **AMINO** TTG TGT GCA -22 Met ATG Ala GCC ATT CGT CAT AAG CAT Ala GCT GTT TIG TTT CTT AND ACG TTC GTG Leu GTT TCT ATT DNA

238

AGA

286

ACG

334

Leu TTA

917

76

AAA

142

ATT

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526	574	622	670	718	166
Trp TGG	Gly	Asp	Phe	120 Lys AAA	Lys
Tyr	Ser	GAA	Thr	Ala GCG	Lys
Ile	70 613 660	Thr	10 10 10	Asp	Ala GCT
Leu	Thr	Lys	Pro CCC	Asp	Asp
CTG	Phe	Val	100 Tyr TAT	Ala GCG	Asp
Lys	Argo	Ser	Ser	Ser	Lys
S C C C C	Asp	Ser	Tyr	Lear	130 Lys AAG
Ser	Pro	Ile	TAT	Hind Lys AAG	Ala
Gln	Val	80 Ser TCC	Glu	CTG	Asp
Gly	GIY	Lec	Glu	Val	Asp
Pro	Ser	Thr	CYS	110 Leu CTG	Lys
Lys	Glu	Phe TTC	Tyr	Lys	Lys
Gln	60 Arg Agg	Asp	Tyr	Thr	Ala
Gln	Ala GCT	Thr	Val	GLY	Ala
Tyr	Ser	<b>G1y</b> GGG	90 Ala GCA	Ala	Asp
Tr TGG	Ala	Ser	Leu	GIY	Lys

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814	862	910	958	1006	1054
or H	Thr	Glu	GAG GAG	Thr	TAT
န	Phe	Leu	AAT Asn AAT	Ser	Val GTG
150 Val L G AAA	Thr	GLY	74C 74C 74C	Ser	230 Ala GCA
วี อาก กับ	Tyr	Gln	AAA Lys AAA	Ser	Ser
Glul	GIY	180 Glu GAA	777 777 777	Lys	Asp
Ala (T. GA(	Ser	Pro	GAT	Asp	Glu GAG
Asp C GC	Ala	Asn	GAT	210 Ala GCA	Ser
Ser J T GA(	Lys	Gln	HP- Asn AAT	Thr	Thr
Gln G TC	160 Cys 1 TGC	Lys	C49VH Gly A GGA A	Leu	Leu
Gln (	Ser TCC	Val	ora CCC	Thr	Ser
Leu G CA	Ile	7 7 7 7 7 7 7	190 Ser TCT	Ala GCC	Asn
Gln G TT	Lys	His	Phe	Lys	CT C
140 Val ( T CA(	Val	Ile	Tyr	Gly	220 Gln CAG
VH I Glu G GT	Ser	Ala	Gly	Lys	Val
Xho Leu C GA	Ala	170 His CAT	I16 ATT	Phe	Tyr
SP CTC	14 1 66 0	A C D	20	r' 0 R 0	ila CC

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1342	Pro	Ser	Gln	GIY	Pro	Lys AAA	Glu	Gln	320 Tyr TAC	44 44 44	Ala	Leu TIG	Tyr	Asn	Lys	AA
1291	Asn	GLY	310 Ser AGT	Tyr	Leu TTA	Lec	Ser	Gln	Ser	Ser	Lys	S S S S S S S S S S S S S S S S S S S	300 Ser AGC	Leu	S. Fr	Th AC
1246	Lys	Glu	GGC	Val	Ser	Val	290 Pro CCT	Leu	Ser	Ser	Pro	Ser	Glu	Ser	42 (5)	Me AT
1198	280 11e ATT	VL Asp GAC	Leu	Asp	Lys	Lys	Ala	ASP	Asp	Lys	270 Lys AAA	Ala	Asp	ASP	n	LY
1150	Ala GCT	Ala GCA	Asp	Lys	260 Lys AAG	Ala GCA	Asp	Asp	Ala GCA	Ser	Leu	Ser	Ser	Val	0 4 0	25 Th AC
1102	Ser TCA	Thr	GLY	CAA	667 667 667	400 400 400	TAC	A18 600	ATG ATG ATG ATG	AAT Asn AAT	7 6 1 C 1 C 1 C 1 C 1 C 1 C 1 C 1 C 1 C 1	NH49J Ser TCC	Arg AGA	Thr	n H	Cy TG

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1390	1438	1486	1534	1582	1630	1678
Asp	360 Ser AGC	TAT Tyr TAT	Leu CTA	Lys	Asp	440 Ala GCT
Pro	ILBATC	TAT Tyr TAT	Lys	Ala	Ser	Lys
Val	Ser	CAG Gln CAG	390 Leu CTG	ASP	Gln	Cya
G13 GGG	Leu	CAG	Val	Asp	Gln	Ser
340 Ser TCT	Thr	13+- Cys 7GT	Leu	Lys	420 Leu TTG	I1e ATT
Glu	Phe	TCDR Tyr TAC	Lys	Lys	Gln	Lys
Arg AGG	Asp	370 49V Tyr TAT	Thr	Ala	Val	Val
Ala GCT	Thr	Val	G13 GGG	Ala	VH Glu GAG	Ser
Ser	GIY	Ala	Ala GCT	400 Asd GAC	CTG	Ala
Ala GCA	Ser	CTe CTG	Gly	Lys	Asp	612 666
Trp	350 614 66A	ASD	Phe	Lys	Lys	#30 Pro CCT
TAC	Ser	Glu	Thr	Ala GCT	Lys	Lys
Ile	G14 GGC	Thr	380 Leu CTC	Asp	Ala GCA	Val
Leu	Thr	Lys	Pro CCC	II Asp GAT	Asp	Leu
330 Leu CTG	Phe	Val	TAT Tyr TAT	47 II Ala GCT	410 Asp Gat	Glu GAG
Lys	Arg	Ser	AGC Ser AGC	Ecol Ser Agc	Lys	Ala GCT

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1966	1918	1870	1822	1774	1726
GAT	520 Tyr TAC	Ser	Ala GCA	Asp	Asn
I AGC	Ala	Thr	Thr	Asn	Gln
Nhe	Met	Leu	Cig	470 G1y GGA	Lys
AAA	Asn	Ser	Thr	Pro	Val
*** TAA	CHO CHO	500 Asn AAC	Ala GCC	Ser	Trp
Ser	NOT CO	Cic	Lys	Phe	His
530 Ser TCC	Arg	Gln	Gly	Tyr	450 Ile ATT
Val	Thr	Val	Lys	Gly	Ala GCA
Thr	A C C C C C C C C C C C C C C C C C C C	Tyr	480 Phe TTC	ILE	His
Val	Phe	Ala	Ark	Tro	Asp
Ser	510 Tyr TAT	Thr	Glu	Glu	Thr
Thr	Val	Ser	Asn	Leu	Phe
Gly	Ala GCA	Ser	Tyr	460 G1y GGC	Thr
Gln	Ser	Ser	Lys	Gln	Tyr
GIY	Asp	490 Lys	Phe	Glu	Gly
1 100 000	Glu	Asp	Asp	Pro	Ser
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JAA	100	GTC S	AAA JP 1-	SAN TCC GTC AAA ACA TCA TCT TAC ATA SOP1- TGT AGT AGA ATG TAT	TCA	TCT AGA	TAC	ATA TAT	AAG	TCA	CH	GGT	TCA CTT GGT GAT CAA GCT AGT	CAA	GCT	2014	
			·		PENPTSEQ2-	rseo:	5 - G	TAT	TTC	AGT	GAA	CCA	CTA	GTT			
CAT	ATC	ATT	GIC	CAT ATC ATT GTC CGG CAA TGG TGT GGG CTT	CAA	TGG	TGT	වච්ච	CTT	TTT	TTG	TTT	TIT TIG III TCT AIC III	ATC	TTT	2062	
3	GAT	CAT	GTG	GAT CAT GTG AAG AAA AAC GGG AAA ATC	AAA	AAC	වුව	AAA	ATC	GGT	CTG	550	GGT CTG CGG GAA AGG ACC	AGG	ACC	2110	
366	TII	TTG	TCG	SGG TTT TTG TCG AAA TCA TAG GCG AAT GGG	TCA	TAG	<b>909</b>	AAT	999	TTG	GAT	TGT	TTG GAT TGT GAC AAA ATT	AAA	ATT	2158	
BamH	HH I	ر ا	-													2165	

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DNA AND AMINO ACID SEQUENCE OF CC49 VL-L-VH-L-VL

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#6	<b>76</b>	142	190	238	286	334	382	#30
ICC	AAA	ATT	TTA	AGA	ACG	Leu	Pro	Lys
SCC	GAA	TAC	CAA	GAA	GAG	Leu TTA	Ser	10 10 10 10
CTT	TCC	GAC	TAC	676 676	AGG	Leu	Gln	Ser
TCA	CCT	AAA	ACG	ATA	<b>9</b> 00	GIY	Ser	Leu
CCA	TTT	TIC	CGT	000 000	AAA TTT	Ala	Met	20 Thr ACT
4 1	TCA	ACT	GAT	GTA	TTC	Ala	Val	Val
TGA A	TCA	AAT	AAA	ACT	ATA TAT	Ala GCA	Ile	Lys
		TAT	CTG	AAC	CAA	Thr	VL Asp GAC	Glu
TTG ACA GCT TAT CAT CGA	GTG GAA ACG AGG	TAC ATA TAT	555		AAT CAA PENPR2-	Pro CCT	Ala GCC	GLY
TAT	GAA	TAC	AGT	PENPR1- AGC CAT	ATC	146 146	Met ATG	Val
GCT	GTG	TCT		TCA		Leu	Nco Ala GCC	Ser
ACA	ဗ္ဗ	AAA TCT	IG ATT TGA TGT TTG	TTT CGT GAT TGT TCA	GTT	Tyr	Pro	Val
TTG	TCC		TGA	GAT	CTG	Lys	Gln	Pro CCH
TGT	TTG	GCA	ATT	CGT	CAT	-22 Met ATG	Ala	Leu
TCA TGT	CAT TTG	GTT GCA TTT	AAG	TTT	CTT CAT	116	Ala GCT	Ser TCC
<u>ي</u>	GTT	ACG	TGI	TTG	GTG	ATT	CTC	Ser

11/20

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## F16. 7E

TG GCC 478	yr Trp AC TGG 526	er Gly GT GGA 574	lu Asp AA GAC 622	hr Phe cg Trc 670	120 12 Lys CG AAA 718	Lys Lys AAA AAG 766
TAC TT	ILO TATE	70 G1y S GGC A	Thr G ACT G	Leu I	Asp A GAT G	Ala L GCT A
ASD	Leu	Thr	Lys	870 000	Asp	ASPGAT
Lys	Leu	Phe	Val	100 Tyr TAT	Ala	ASP
Gln	Lys	Argo	Ser	Ser	Ser	Lys
Asn AAT	GCT O	Asp	Ser	Tyr	Lectr	130 Lys AAG
Gly	Ser	Pro	Ile	Tyr	Hind Lys AAG	Ala GCT VL(-
Ser	Gln	Val	Ser TCC	Gln	CHO	ASP GAC TMN
Tyr	G13 GGG	919 999	Leu	Gln	Val	Asp GAT CTA
Leu TTA	Pro	Ser	Thr	Cys	Leuc	Lys AAG TTC
Leu	Lys AAA	Glu	Phe	Tyr	Lys	Lys AAG TTC
Ser	Gln	Arg AGG	Asp	Tyr	Thr	A1a 666 666
Gln	Gln	Ala	Thr	Val	GIY	Ala GCT CGA
Ser	Tyr	Ser	GIY	90 Ala GCA	Ala	Asp GAT CTA
Ser	Trp	Ala	Ser	Cito	GIY	Lys

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1102	1054	1006	958	910	862	814
Ser	TAT	Thr	200 GAG GAG	Glu	Thr	Pro
Thr	Val	Ser	AAT Asn AAT	Leu	Phe	Lys
G G G G A A	230 Ala GCA	Ser	TAC	Gly	Thr	Val
CAA Gln CAA	Ser	Ser	AAA Lys Aaa	Gln CAG	Tyr	Leu
667 617 667	Asp	Lys	TTT Phe TTT	180 G1u GAA	G17 GG24	Glu
166 170 166	Glu	ASD	GAT GAT	Pro	Ser	Ala GCT
TAC Tyr TAC	Ser	210 Ala GCA	GAT	Asn	Ala	Asp
GCC Ala GCC	Thr	Thr	VHP- Asn AAT	Gln	Lys	Ser
240 ATG Met ATG	Leu	CTO	0 4 9 0 4 9 0 4 9	Lys	160 Cys TGC	Gln
AAT AAT	Ser	Thr		Val	Ser	Glu
J- G CTG	Asn	Ala	190 Ser TCT	77 700 700	Ile ATT	Leu
VH49. Ser TCC	Lec	Lys	Phe	HIS	Lys	Gln
Arg	220 Gln CAG	GLY	TAT	I1e ATT	Val	Val
Thr	Val	Lys	GIY	Ala	Ser	GIU
Cys	Tyr	Phe	ILE	170 H13	Ala GCT	CHC
Phe	Ala GCC	Arg	100	Asp	<b>61y</b> 666	Asp

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1150	1198	1246	1294	1342	1390	1438	1486
a H	280 Val GTT	Val	ILE	Tyr	Gly	360 Gln CAG	Arg
18 A CA G	VH Glu GAG	Ser	Ala GCA	GGA	Lys	Val	Thr
SP A	Leu	Ala GCT	310 His CAT	ILGATT	Phe	Tyr	Cys
Lys A	Asp	Gly	ASP	44	Arg	Ala GCC	Phe
84 60 AG 40	Lys	Pro	Thr	GAA	340 G1u GAG	Thr	Tyr
CA	Lys	Lys	Phe	CTO	Asn	Ser	Val
ASP A	Ala	290 Val GTG	Thr	GIY	Tyr	Ser TOC	370 Ala GCA
SP A	ASP	Leu	Tyr	Gln	Lys AAA	Ser	Ser
Lla A	Asp	Glu	GIY	320 61u 6AA	Phe	Lys	Asp
Ser A	Lys	Ala GCT	Ser	8 6 6 7	ASP	Asp	Glu
Leu S CTA A	270 Lys AAA	ASP	Ala GCT	Asn	Asp	350 Ala GCA	Ser
Ser I	Ala	Ser	Lys	Glu	Asn	Thr	Thr
Ser	Asp	Gln	300 Cy 8	Lys	Gly	Leu	Leu
Val GTC	Asp	Gln	Ser	Val	Pro	Thr	Ser
250 Thr ACC	Lys	Leu	Ile	Trp	330 Ser TCT	Ala	ABD
/al	Lys	Gln	Lys	HIS	Phe	Lys	Leu

14/20

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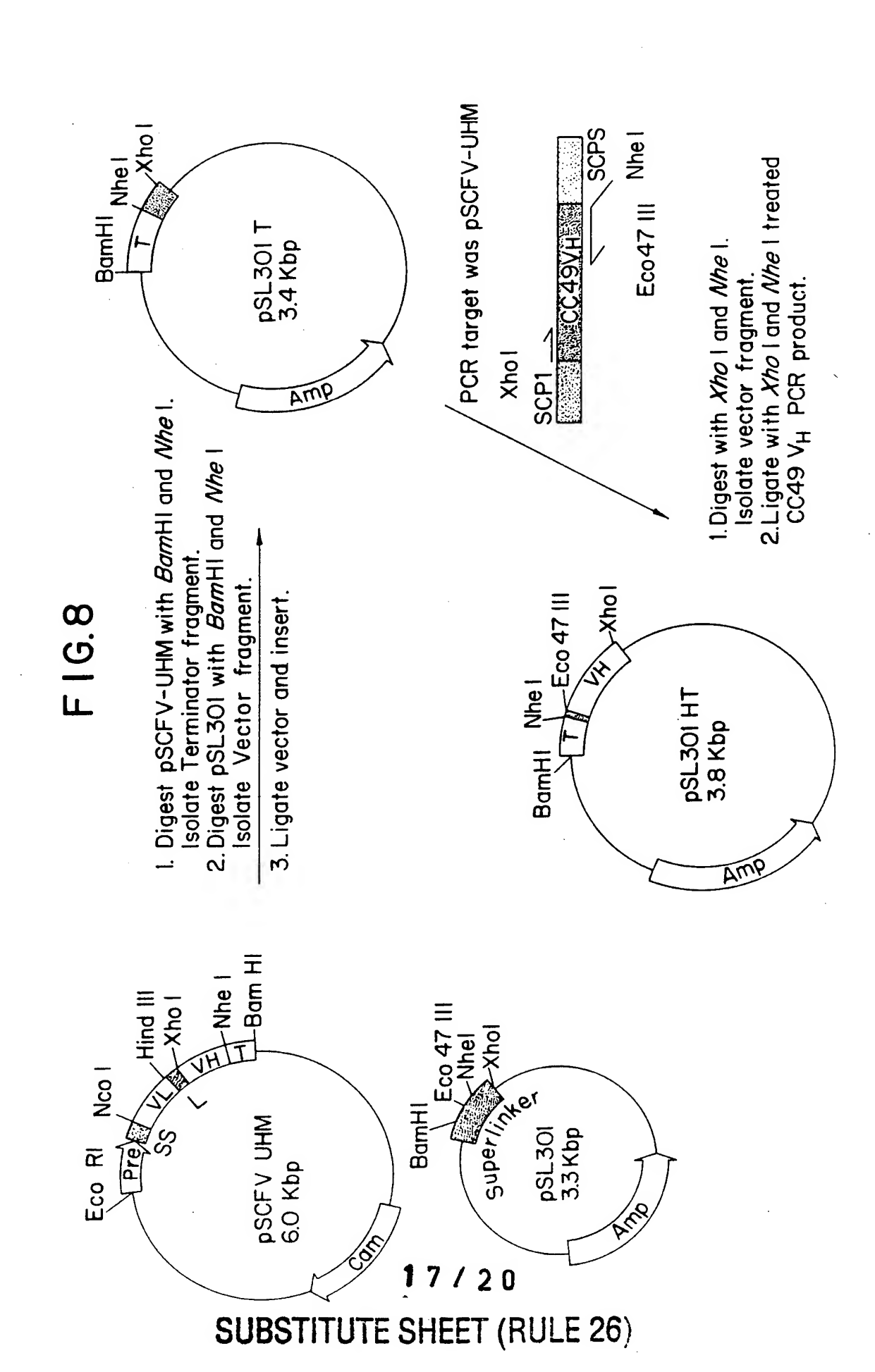
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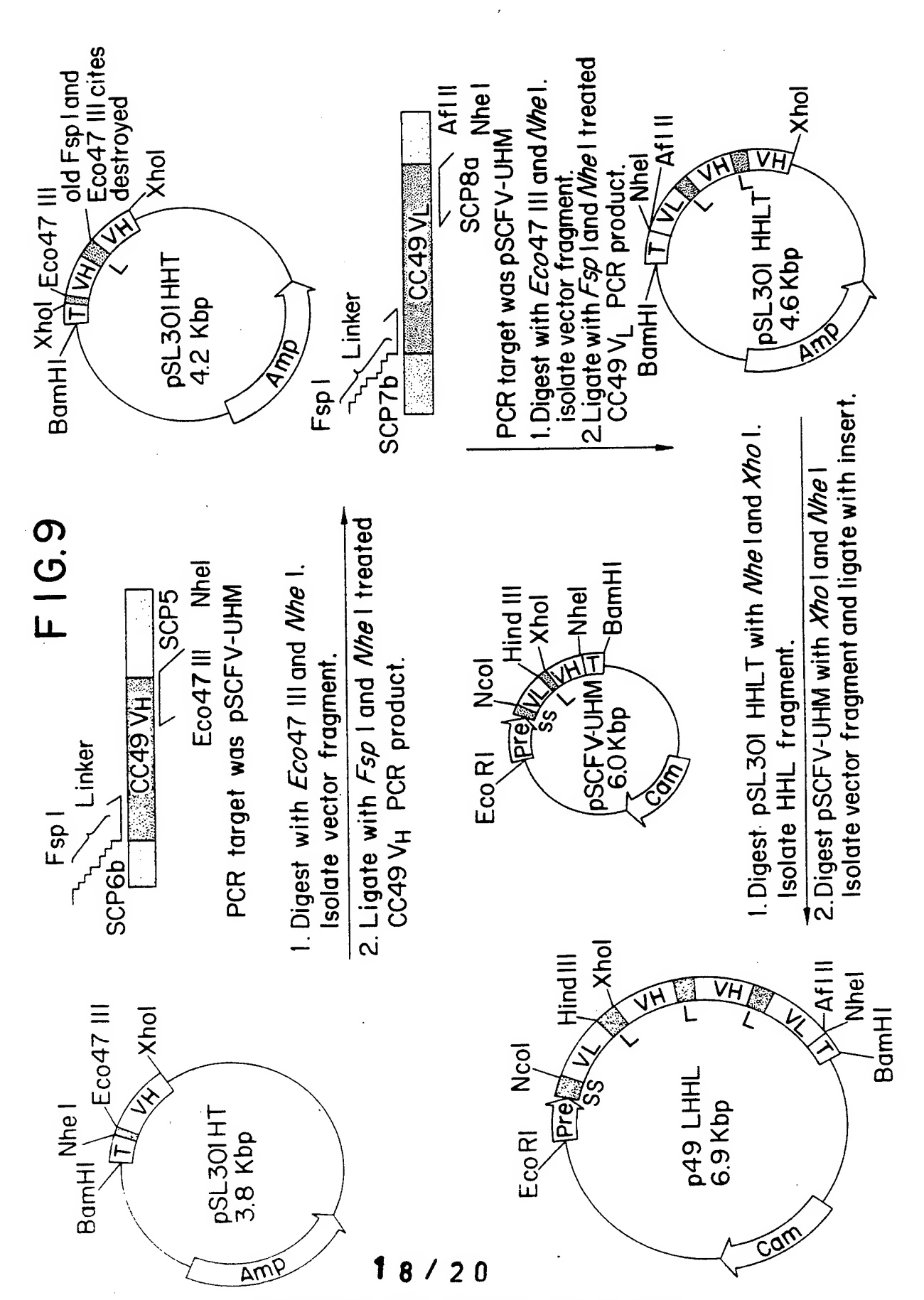
1870	1918	1966	2014		2062	2110	2158	2165
Thr	520 Leu CTC	GAT	GCT		TTT	ACC	ATT	
Lys	Pro CCC	IAGC	CAA	GTT	ATC	AGG	AAA	
Val	Tyr	Nhe	GAT	CTA	TCT	GAA	GAC	
Ser	Ser	AAA	GGT	CCA	TTT	<b>99</b> 0	TGT	
Ser AGC	Tyr	*** TAA	CTT	GAA	TIG	CTG	GAT	
Ile	Tyr	II Lys AAG	TCA	AGT	TTT	GGT	TTG	
Ser	Gln	Afri CHT CHT	AAG	TTC	CTT	ATC	ອອອ	
Leu	Gln	Val	_ <b>[</b> •	TAT	999	AAA	AAT	
Thr	Cys	Leu	TAC	A A1G	TGT	000	ອວອ	
Phe TTC	Tyr	Lys	TCT	A S E	TGG	AAC	TAG	
Asp	510 Tyr TAT	Thr	TCA	ンス	CAA	AAA	TCA	
Thr	Val	GIJY	ACA	5	<del>ງ</del>	AAG	AAA	
Gly	Ala GCA	Ala GCT	AA	-L 40	GIC	GTG	TCG	•
Ser	Lea	Gly	GTC	Š	ATT	CAT	TTG	C-3
490 Gly GGA	Asp	Phe	TCC		ATC	GAT	TTT	Bamh I GG ATC
Ser	G1u GAA	Thr	GAA		CAT	AAA	ງວງ	Bay

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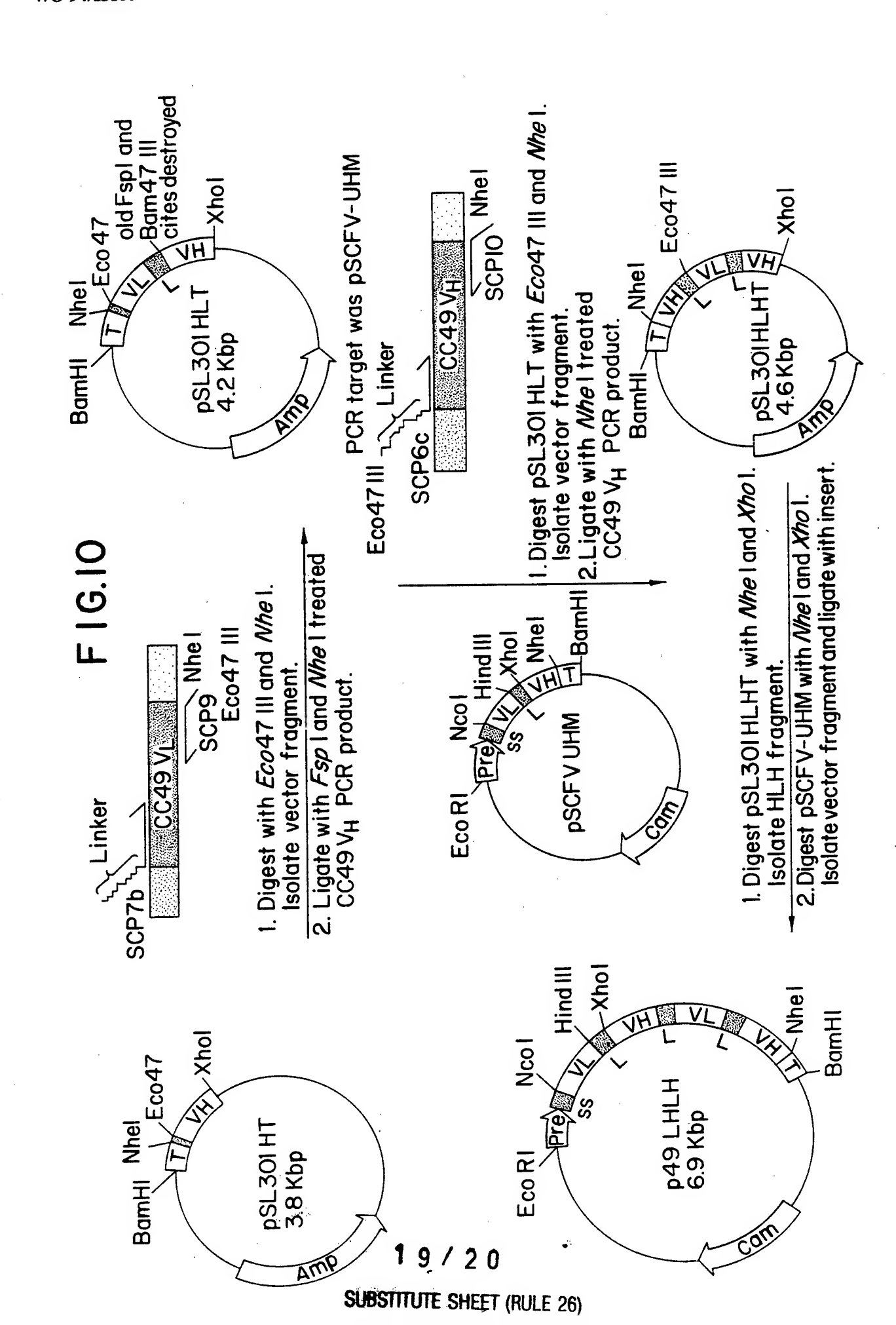
1534	1582	1630	1678	1726	1774	1822
Ser	Asp	Gln	Ser AGC	TAC TAC	ILEATT	GIY
Val	ASP	Ser	Leu	Asn AAC R2(-)	Cig	Thr
390 Thr ACC	Lys	Met	Thr	Lys AAG 9LF	470 Leu CTG	Phe
Val	Lys	Val	Val	Gla	Lys	Arg
Ser	Ala	420 11e ATT	Lys	Asn	Pro	Asp
Thr	Ala	VL Asp Gac	Glu	GIY	Ser	Pro
Gly	Asp	Leu	GIY	450 Ser AGT	Gln	Val
Gln	Lys	Asp	Val	TAT	613 666	Gly
Gly	400 Lys AAG	Lys	Ser	Leu	Pro	480 Ser TCT
7rp 766	Ala	Lys	Val	Leu	Lys	Glu
TAC	ASP	Ala	430 Pro	Ser	Gluca	Arg
Ala GCC	Asp	ASP	Leu	Gln	GID GAG GTC	Ala
380 Met ATG	Ala	Asp	Ser	Ser	460 Tyr Tac Atg	Ser
Asn	Ser	Lys	Ser	Ser	Tro TGG ACC	Ala GCA
Leu	Leu	410 Lys	Pro	Lys	A18 GCC CGG	7 7 7 7 7 7 7
Ser	Ser	Ala GCC	Ser	Cya	Leu TTG AAC	Tyr

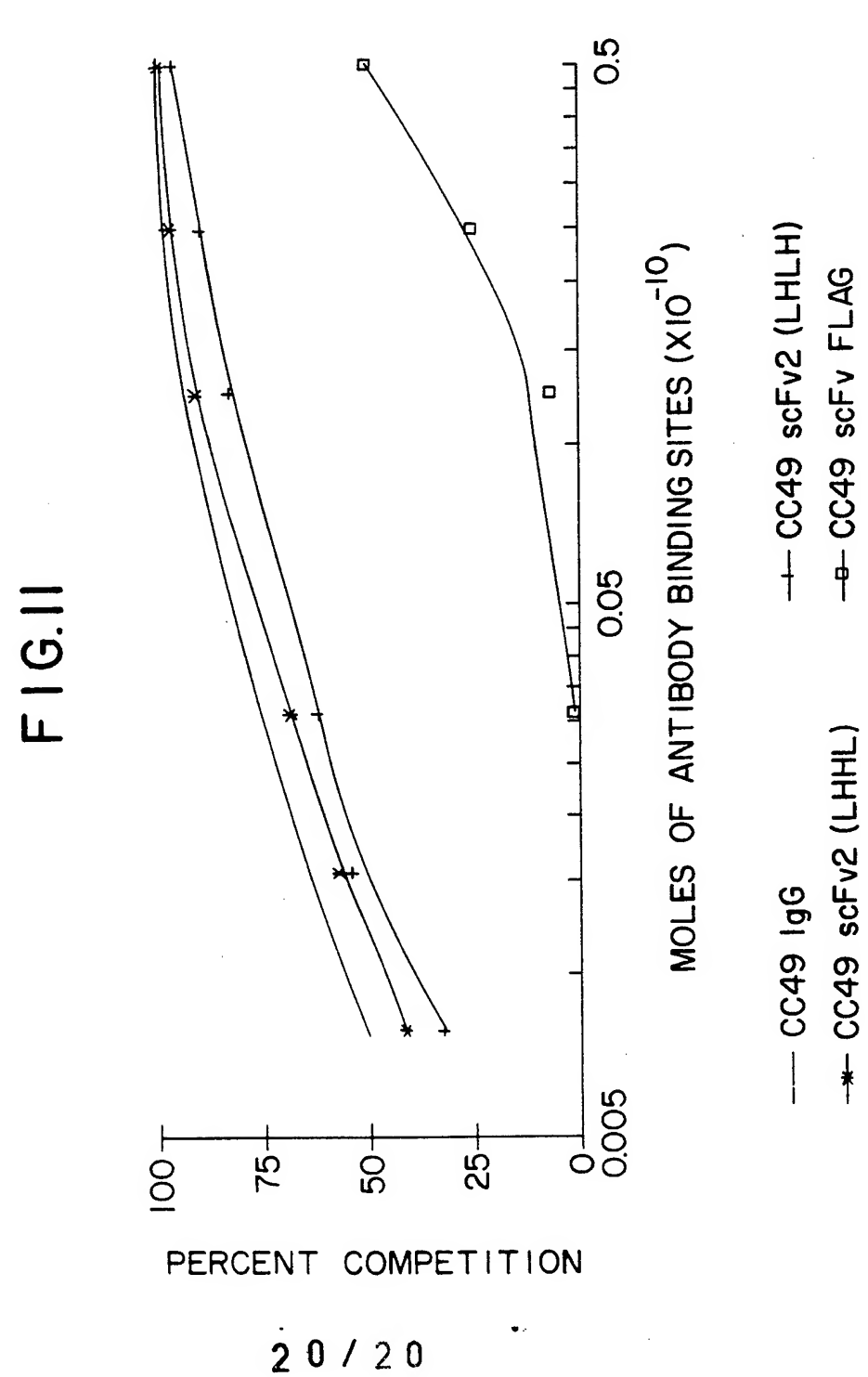
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PCT/US 93/12039 A. CLASSIFICATION OF SUBJECT MATTER IPC 5 C12N15/13 C07K15/28 A61K39/395 C12N15/62 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) C12N C07K IPC 5 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category \* 1,5 WO, A, 91 19739 (CELLTECH LIMITED) 26 X December 1991 2-4,6Y see example 1 3,6 Y CANCER RESEARCH vol. 52, no. 12 , 15 June 1992 , PHILADELPHIA, PA, USA pages 3402 - 3408 T.YOKATA ET AL. 'Rapid tumour penetration of a single-chain Fv and comparison with other immunoglobulin forms' see page 3403, column 1, paragraph 4 Patent family members are listed in annex. Further documents are listed in the continuation of box C. • Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention filing date cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another "Y" document of particular relevance; the claimed invention citation or other special reason (as specified) cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means in the art. document published prior to the international filing date but "&" document member of the same patent family later than the priority date claimed Date of mailing of the international search report Date of the actual completion of the international search 27 -04- 1994 25 March 1994 Authorized officer Name and mailing address of the ISA

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